Organic Reactions

Organic Reactions

VOLUME VII

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PREFACE TO THE SERIES

In the course of nearly every program of research in organic chemistry the investigator finds it necessary to use several of the better-known synthetic reactions. To discover the optimum conditions for the application of even the most familiar one to a compound not previously subjected to the reaction often requires an extensive search of the literature; even then a series of experiments may be necessary. When the results of the investigation are published, the synthesis, which may have required months of work, is usually described without comment. The background of knowledge and experience gained in the literature search and experimentation is thus lost to those who subsequently have occasion to apply the general method. The student of preparative organic chemistry faces similar difficulties. The textbooks and laboratory manuals furnish numerous examples of the application of various syntheses, but only rarely do they convey an accurate conception of the scope and usefulness of the processes.

For many years American organic chemists have discussed these problems. The plan of compiling critical discussions of the more important reactions thus was evolved. The volumes of Organic Reactions are collections of chapters each devoted to a single reaction, or a definite phase of a reaction, of wide applicability. The authors have had experience with the processes surveyed. The subjects are presented from the preparative viewpoint, and particular attention is given to limitations, interfering influences, effects of structure, and the selection of experimental techniques. Each chapter includes several detailed procedures illustrating the significant modifications of the method. Most of these procedures have been found satisfactory by the author or one of the editors, but unlike those in Organic Syntheses they have not been subjected to careful testing in two or more laboratories. When all known examples of the reaction are not mentioned in the text, tables are given to list compounds which have been prepared by or subjected to the reaction. Every effort has been made to include in the tables all such compounds and references; however, because of the very nature of the reactions discussed and their frequent use as one of the several steps of syntheses in which not all of the intermediates have been isolated, some instances may well have been missed. Nevertheless, the investigator will he at !!

to use the tables and their accompanying bibliographies in place of most or all of the literature search so often required.

Because of the systematic arrangement of the material in the chapters and the entries in the tables, users of the books will be able to find information desired by reference to the table of contents of the appropriate chapter. In the interest of economy the entries in the indices have been kept to a minimum, and, in particular, the compounds listed in the tables are not repeated in the indices.

The success of this publication, which will appear periodically, depends upon the cooperation of organic chemists and their willingness to devote time and effort to the preparation of the chapters. They have manifested their interest already by the almost unanimous acceptance of invitations to contribute to the work. The editors will welcome their continued interest and their suggestions for improvements in *Organic Reactions*.

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CHAPTER 1

THE PECHMANN REACTION

Suresh Sethna * and Ragini Phadke

Royal Institute of Science, Bombay

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INTRODUCTION

H. v. Pechmann found that coumarin derivatives are formed when malic acid 1 or β -ketonic esters 2 are condensed with phenols in the presence of concentrated sulfuric acid. This reaction, which is commonly known as the Pechmann reaction, has found extensive application.

HO
$$\begin{array}{c}
\text{OH} + \text{CO}_2\text{H} \\
\text{CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}
\end{array}$$

$$\begin{array}{c}
\text{H}_2\text{SO}_4 \\
\text{HOCCH}_3
\end{array}$$

$$\begin{array}{c}
\text{H}_2\text{SO}_4 \\
\text{HOCCH}_3
\end{array}$$

$$\begin{array}{c}
\text{H}_2\text{SO}_4 \\
\text{HOCCH}_3
\end{array}$$

$$\begin{array}{c}
\text{CH}_3
\end{array}$$

Simonis and his co-workers 2.4.5 used phosphorus pentoxide as the condensing agent in place of sulfuric acid and demonstrated that with the same reactants chromones rather than coumarins resulted. It was

$$OH + C_2H_5O_2C \longrightarrow CO$$

$$C_2H_5O_2C \longrightarrow CO$$

shown later, however, that chromones were not always the reaction products. The condensation of a phenol and β -ketonic ester in the presence of phosphorus pentoxide is sometimes called the Simonis reaction,

¹ v. Pechmann, Ber., 17, 929 (1884).

Pechmann and Duisberg, Ber., 16, 2119 (1883).
 Fewchek and Simonis, Ber., 46, 2014 (1913).

^{*} Simonis and Lehmann, Ber., 47, 692 (1914).

³ Simonia and Remmert, Ber., 47, 2229 (1914).

but it is actually merely a variation of the Pechmann reaction and will be so considered in this chapter. Other condensing agents that have been used are phosphorus oxychloride, phosphoric acid, zinc chloride, aluminum chloride, hydrogen chloride, ferric chloride, stannic chloride, titanic chloride, sodium acetate, sodium ethoxide, and boric anhydride.

By condensing appropriately substituted phenols and β -ketonic esters, coumarins can be synthesized with substituents either in the benzene nucleus or in the heterocyclic ring or in both. These compounds can then be used for the preparation of other products like coumarino- α -pyrones, coumarino- γ -pyrones, furocoumarins, chromenes, coumarones, and 2-acylresorcinols.⁶ The Pechmann reaction has also been employed in the syntheses of several naturally occurring coumarins ^{6,7} and in the investigations of natural products like rotenone ⁸ and cannabinol.^{9,10}

The course of this reaction depends on all of the three factors: the nature of the phenol, the nature of the β -ketonic ester, and the condensing agent.

MECHANISMS OF THE REACTIONS

Condensation of Malic Acid with Phenols. The condensation of malic acid with phenols takes place according to Pechmann ¹ in three stages. The malic acid is first converted into malonaldehydic acid and formic acid, which is decomposed into water and carbon monoxide.

$$\text{HO}_2\text{CCH}(\text{OH})\text{CH}_2\text{CO}_2\text{H} \rightarrow \text{HCO}_2\text{H} + \text{CHOCH}_2\text{CO}_2\text{H}$$

In the second stage, the union of the aldehyde with the phenol results in the formation of an unstable addition product. Two molecules of water are then eliminated, and the coumarin derivative is formed. Malonaldehydic acid contains a carbonyl group in the β position and resembles ethyl acetoacetate in its reaction with a phenol to give a coumarin.

⁶ Sethna and Shah, Chem. Revs., 36, 30 (1945).

⁷ Spath, Ber., 70A, 83 (1937).

⁸ Bridge, Crocker, Cubin, and Robertson, J. Chem. Soc., 1937, 1530.

⁹ Ghosh, Todd, and Wilkinson, J. Chem. Soc., 1940, 1121.

¹⁰ Adams and Baker, J. Am. Chem. Soc., 62, 2405 (1940).

Condensation of β-Ketonic Esters with Phenols. To explain the formation of coumarins from β-ketonic esters and phenols, Pechmann and Duisberg 2 suggested that the reactive hydrogen of the phenol in the ortho position to the hydroxyl group adds to the carbonyl of the β-ketonic ester to give an intermediate hydroxy ester (I). Ring elosure may then take place with the elimination of a molecule of water and one of ethanol.

Ahmad and Desai 11 have pointed out that the effectiveness of such condensations depends on the reactivity of the hydrogen in the ortho-

$$OH + COCH_{2}CO_{2}C_{2}II_{5}$$

$$CH_{3}$$

$$CH_{$$

position to the hydroxyl group and on the substituents in the β-ketonic ester. The feeble tendency of phenol itself to condense is enhanced by the presence of electron-donating groups such as CH₃, OH, OCH₃, NH₂, NHCH₃, N(CH₃)₂, and halogens in the *meta* position to the hydroxyl group but is depressed or almost eliminated by electron-attracting groups such as NO₂, SO₃H, CO₂H, CO₂CH₃, COCH₃, CN, and CHO in the same position.¹² Since no intermediates have been isolated this course for the reaction is purely speculative.

A slightly different view has been advanced by Robertson and his co-workers. They observed that 2-methoxy-\$\beta\$,4-dimethylcinnamic acid was converted into 4,7-dimethylcoumarin in the presence of 86% sulfuric acid and, further, that m-tolyl methyl ether and the dimethyl ether of resorcinol gave rise to 4,7-dimethylcoumarin and 7-methoxy-4-methylcoumarin, respectively. From this experimental evidence they conclude that the cinnamic acid derivative (II) is formed as an intermediate product.

¹³ Robertson, Waters, and Jones, J. Chem. Soc., 1932, 1681.

Ahmad and Desai, Proc. Indian Acad. Sci., 6A, 6 (1937) [C. A., 32, 559 (1938)].
 Desai and Ekhlas, Proc. Indian Acad. Sci., 8A, 567 (1938) [C. A., 33, 3356 (1939)].

$$\begin{array}{c} \text{H}_3\text{C} \\ \text{C} = \text{CHCO}_2\text{H} \\ \text{CH}_3 \end{array} \xrightarrow{\text{H}_3\text{C}} \begin{array}{c} \text{O}_{\text{CO}} \\ \text{CH}_3 \end{array}$$

Two different mechanisms for chromone formation have been proposed. Robertson and his co-workers suggest that the first stage in the reaction results in a phenoxy acid (or its ester) by the interaction of the enolic form of the ester and phenol with the removal of a molecule of water. The phenoxy derivative then undergoes ring closure to a chromone. In support of this mechanism they cite the synthesis of

$$\begin{array}{c} \text{OH} & \text{HOCCH}_3 \\ & \text{C}_2\text{H}_5\text{O}_2\text{C} \\ & \text{C}_2\text{H}_5\text{O}_2\text{C} \\ & \text{C}_2\text{H}_5\text{O}_2\text{C} \\ & \text{CO} \\ \end{array}$$

chromones from phenoxyfumaric acid and β -phenoxyfinnamic acid by

According to Ahmad and Desai, in the formation of chromones, the reactive hydrogen of the phenolic hydroxyl reacts with the exposed, the reactive hydrogen of the phonon street of the acid (III). This results in the β -ketonic ester to give an arylester of the acid (III). This results in the phonon is β-ketonic ester to give an ary cooperation is based on the evidence that only those phenois that do not contain a reactive hydrogen ortho to the hydroxyl group give chromones in the presence of phosphorus pentoxide as condensing agent. The series exerthen undergoes an isomeric change analogous to the Fire toleration (IV) which is determined to the property toleration. forming an o-hydroxybenzoylacetone (IV) which is dehive to the transfer to the chromone derivative (V). They assume the transformed to the Schapherg and M. possible in view of the work of Schönberg and Musican to be rearrangements with phosphorus pentoxide. They suggested that the state of the stat the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of phosphorus pentoxide is to facilitate the specific action of the specific

¹⁴ Ruhemann and co-workers, J. Chem. Soc., 77, 984, 1119 (1906); 73, 577, 1117

of III or IV or both since the conversion of IV into V may be accomplished with the help of any dehydrating agent. The formation of the

$$\begin{array}{c}
\text{OH} & \text{CO}_2\text{C}_2\text{H}_5 \\
\text{COCH}_3 & \text{COCH}_3
\end{array}$$

$$\begin{array}{c}
\text{OH} \\
\text{COCH}_2\text{COCH}_3
\end{array}$$

$$\begin{array}{c}
\text{OH} \\
\text{COCH}_2\text{COCH}_3
\end{array}$$

$$\begin{array}{c}
\text{OH} \\
\text{COCH}_2\text{COCH}_3
\end{array}$$

intermediate diketone IV in the syntheses of chromones by the Kostanecki acylation of o-hydroxyketones has been proved by Baker. 16

Formation of 5-Hydroxycoumarin Derivatives in Presence of Anhydrous Aluminum Chloride. The formation of 5-hydroxycoumarin derivatives in the condensations of resacetophenone, 4-nitroresorcinol, and methyl β-resorcylate in preference to the 7-hydroxycoumarin derivatives is obviously due to the greater reactivity of the usually inaccessible 2-position of the resorcinol nucleus in these compounds. Shah and Shah ¹⁷ have explained this on the basis of chelation between the hydroxyl group and the *ortho*-substituted group, thus fixing the double bonds. ^{18, 19, 20} The point of attack is consequently the carbon atom joined by a double bond to that bearing the other hydroxyl group; resacetophenone and ethyl acetoacetate condense with the formation of 5-hydroxy-6-acetyl-4-methylcoumarin. The formation of a 5-hydroxy-coumarin from methyl β-resorcylate and 4-nitroresorcinol in the presence of aluminum chloride can be explained similarly.

$$CH_3\overset{0}{\underset{\parallel}{\text{C}}}OH + CH_3COCH_2CO_2C_2H_5 \longrightarrow CH_3\overset{0}{\underset{\parallel}{\text{C}}}OH CH_3$$

Baker 19 believes that aluminum chloride may prevent chelation; but, since 5-hydroxycoumarins are formed mainly or exclusively in good yields in the above condensations, it appears that this reagent not only fails to prevent chelation but may even promote it, for other condensing

¹⁸ Baker, J. Chem. Soc., 1933, 1381.

¹⁷ Shah and Shah, J. Chem. Soc., 1938, 1424.

¹⁸ Mills and Nixon, J. Chem. Soc., 1930, 2510.

¹⁸ Baker, J. Chem. Soc., 1934, 1684.

²⁰ Baker and Lothian, J. Chem. Soc., 1935, 628.

agents generally produce derivatives of 7-hydroxycoumarin. This view also finds support in the work on the formylation of methyl β -resorcylate 21 and 4-acylresorcinols; 22,23 the Gattermann reaction in the presence of anhydrous aluminum chloride in dry ether leads to formylation in the 2 position, in the case of resacetophenone yielding 2-formyl-resacetophenone.

SCOPE AND LIMITATIONS

The reactivity of the various simple and substituted phenols and β -ketonic esters in the Pechmann reaction, with sulfuric acid as the condensing agent, will be discussed first, and the role of the condensing agents second.

Reactivity of Phenols. It is found that, of the simple mono-, di-, and tri-hydric phenols, resorcinol is the most reactive, and it condenses with many substituted and cyclic β -ketonic esters. Almost equal in reactivity are phloroglucinol, α -naphthol, and pyrogallol. Phenol, quinol, and β -naphthol, however, usually give low yields of products. Phenol, for example, gives only about a 3% yield of 4-methylcoumarin on condensation with ethyl acetoacetate in the presence of sulfuric acid,²⁴ and it does not condense at all with many other β -ketonic esters. Catechol does not condense even with ethyl acetoacetate.

Among the substituted phenols it is found that the reactivity depends both on the nature and on the position of the substituent in the phenol. Alkyl groups in general have very little inhibiting effect in the Pechmann reaction; halogens exert somewhat more. When substituents like the nitro and the carboxyl groups are present, the reactions may not take place at all. 25,26 This is exemplified by the non-reactivity of o-, m-, or p-nitrophenol and simple phenol carboxylic acids with ethyl acetoacetate and other β -ketonic esters. The rate and degree to which a coumarin is produced depend, however, on the position of the substituent. m-Cresol condenses very readily with ethyl acetoacetate and a number of other β -ketonic esters, 27,28 p-cresol less readily, 2,28 and o-cresol not at all, even with ethyl acetoacetate. 29 m- and p-Chlorophenols react with ethyl acetoacetate, but o-chlorophenol does not react. 25 m-Dimethylaminophenol condenses with acetonedicarboxylic acid, but the ortho and para

²¹ Shah and Laiwala, J. Chem. Soc., 1938, 1828.

²² Shah and Shah, J. Chem. Soc., 1939, 132.

²³ Shah and Shah, J. Chem. Soc., 1940, 245.

²⁴ Pechmann and Kraft, Ber., 34, 421 (1901).

²⁵ Clayton, J. Chem. Soc., 93, 2016 (1908).

²⁵ Dey, J. Chem. Soc., 107, 1606 (1915).

²⁷ Fries and Klostermann, Ber., 39, 871 (1906).

²⁸ Fries and Klostermann, Ann., 362, 1 (1908).

²⁹ Chakravarti, J. Indian Chem. Soc., 9, 31 (1932).

compounds are inert.26 Thus in many monohydric phenols a substituent in the ortho position has the maximum inhibiting effect, less if the same substituent is in the para position, and least when it is in the mela

position.

The influence of substituents in the resorcinol nucleus on the Pechmann reaction has been investigated. In molecules where substituents in the 4 position cause the reaction to take place with some difficulty, the same substituents in the 2 position have less effect. Resorcinols with alkyl groups in the 2 or 4 position react as readily as résorcinol. Even 4-hexadecylresorcinol condenses smoothly with ethyl acctoacetate in the presence of sulfuric acid.30 Alkyl groups in the 5 position change the course of the reaction, and, instead of the 7-hydroxycoumarin derivatives, the 5-hydroxy isomers are obtained; an exception is in the condensation with malic acid. Thus orcinol 26,31-35 and other 5-alkylresorcinols $^{36-38}$ with ethyl acetoacctate and other β -ketonic esters give 5-hydroxycoumarin derivatives. Orcinol with malic acid gives a 7-hydroxycoumarin.39,40,*

$$+ CH_{3}COCH_{2}CO_{2}C_{2}H_{5} \xrightarrow{H_{2}SO_{4}} H_{3}C \xrightarrow{O} CO$$

$$+ HO \xrightarrow{2} OH \qquad HO CH_{3}$$

$$+ HO_{2}CCH_{2}CHOHCO_{2}H \xrightarrow{H_{2}SO_{4}} HO \xrightarrow{CH_{3}} CH$$

4-Chlororesorcinol condenses smoothly with a number of β -ketonic esters like ethyl a-alkylacetoacetates, ethyl benzoylacetate, and diethyl

- 30 Chudgar and Shah, J. Univ. Bombay, 13, Pt. 3, 18 (1944) [C. A., 39, 4078 (1945)]. 31 Krishnaswamy, Rao, and Seshadri, Proc. Indian Acad. Sci., 19A, 5 (1944) [C. A., 39, 1153 (1945)].
 - 32 Pechmann and Hancke, Ber., 34, 354 (1901).
 - 33 Chakravarti, J. Indian Chem. Soc., 8, 407 (1931).

34 Shah and Shah, J. Indian Chem. Soc., 19, 481 (1942).

- 35 Kotwani, Sethna, and Advani, Proc. Indian Acad. Sci., 15A, 441 (1942) [C. A., 37, 624 (1943)].
 - ²⁶ Russell, Todd, Wilkinson, Macdonald, and Woolfe, J. Chem. Soc., 1941, 169.
 - 37 Russell, Todd, Wilkinson, Macdonald, and Woolfe, J. Chem. Soc., 1941, 826. 33 Adams, Loewe, Jelinek, and Wolff, J. Am. Chem. Soc., 63, 1971 (1941).
 - ³⁹ Pechmann and Welsh, Ber., 17, 1646 (1884).
 - 40 Sastry, J. Indian Chem. Soc., 19, 403 (1942).
 - *7-Hydroxy-4,5-dimethylcoumarin, which cannot be obtained by the direct condensation of orcinol with ethyl acetoacetate, has been prepared by the decarboxylation of 7-hydroxy-4,5-dimethylcoumarin-8-carboxylic acid formed by the condensation of p-orsellinic acid with ethyl acetoacetate. Sethna and Shah, J. Indian Chem. Soc., 17, 211 (1940).

.. . + 53

The capacity of hydroquinone to undergo the Pechmann reaction is not great. When a chlorine atom is present in the hydroquinone the reaction takes place even less readily, and the presence of a bromine atom or acetyl group prevents the reaction completely. On the other hand, greater reactivity is observed when a methyl or ethyl group is substituted in the hydroquinone. 2-Methyl- and 2-ethyl-hydroquinone form commarins with ethyl benzoylacetate and ethyl α -alkylacetoacetates; but quinacetophenone and 2-bromohydroquinone do not condense even with ethyl acetoacetate, and 2-chlorohydroquinone reacts with difficulty. Hydroquinone, its 2-chloro- and 2-bromo-derivative, and quinacetophenone do not condense with ethyl benzoylacetate.56

Of the trihydroxy compounds, 4-ethylpyrogallol and ethyl pyrogallolcarboxylate condense readily with ethyl acetoacetate, ethyl a-alkylacetoacetates, and ethyl benzoylacetate. Gallic acid, its methyl and ethyl esters, pyrogallolearboxylic acid, and gallacetophenone do not undergo the coumarin condensation with these same β-ketonic esters. 57

Phloroglucinol and many of its derivatives, methylphloroglucinol,58 dimethylphloroglucinol,58 methyl phloroglucinolcarboxylate,59 phloroacetophenone, and phlorobenzophenone give coumarins with ethyl acetoacetate. The reaction with other \beta-ketonic esters has not been studied.

1,2,4-Triacetoxybenzene and ethyl acetoacetate in the presence of 75% sulfuric acid condense to give 6,7-dihydroxy-4-methylcoumarin.60 No condensation of a substituted 1,2,4-trihydroxybenzene with a B-ketonic ester has been reported.

α-Naphthol derivatives with chlorine or bromine in the 4 position react with ethyl α-alkylacetoacetates and other β-ketonic esters like ethyl benzoylacetate, diethyl acetonedicarboxylate, and diethyl acetosuccinate. 4-Bromo-α-naphthol appears to be less reactive than 4-chloro-α-naphthol. In the condensation of 4-acetyl-, 4-propionyl-, and 4-butyryl- α -naphthol with β -ketonic esters, the acyl group is eliminated.12 Substituted \(\beta\)-naphthols have not been studied.

Attempts to condense cyclohexanol and dimethyl dihydroresorcinol with acetonedicarboxylic acid did not succeed.26

Certain miscellaneous compounds not included in the previous discussion have been condensed with malic acid and \beta-ketonic esters in the presence of sulfuric acid. Resorcinol and other polyhydroxyphenols

u Desai and Mavani, Proc. Indian Acad. Sci., 15A, 11 (1942) [C. A., 36, 6151 (1942)]. ¹ Devai and Mavani, Proc. Indian Acad. Sci., 15A, 1 (1942) [C. A., 36, 6150 (1942)].

Fujise and Maruyama, J. Chem. Soc. Japan, 55, 1013 (1934) [C. A., 29, 4008 (1935)]. D Sethna, J. Unic. Bombay, 9 (pt. 3), 104 (1940) [C. A., 35, 6948 (1941)].

⁴¹ Vliet, Org. Syntheses, 4, 45 (1924).

⁴ Chakravarti and Bagchi, J. Indian Chem. Soc., 13, 649 (1936).

will not react satisfactorily with two molecules of ethyl acetoacetate or malic acid simultaneously, but the pure hydroxycoumarins formed by the condensation of one molecule of ethyl acetoacetate or malic acid will react with a second molecule of ethyl acetoacetate or malic acid to produce coumarino- α -pyrones. ^{62, 63} The condensation of hydroxycoumarins with malic acid takes place more readily than with ethyl acetoacetate, though the condensation of many simpler aromatic hydroxy compounds with malic acid is more difficult than with ethyl acetoacetate. The dihydroxycoumarins derived from pyrogallol and ethyl acetoacetate will react with malic acid ⁶³ but not with ethyl acetoacetate.

Hydroxychromones do not undergo condensation with malic acid.⁶⁴
Hydroxythiophene derivatives react with β-ketonic esters to yield thiocoumarin derivatives.^{65,66}

Reactivity of Malic, Maleic, and Fumaric Acids. The condensation of malic acid with phenols leads to coumarins which are unsubstituted in the pyrone ring. This procedure is therefore an alternative method for the synthesis of coumarins that are difficult to obtain by Perkin's method from o-hydroxy aromatic aldehydes. There are, however, limitations in the preparation of coumarins by this method: malic acid does not condense with many substituted phenols, and, when it does condense, the yields are often low and tarry products are obtained. Malic acid condenses only in the presence of sulfuric acid; other condensing agents fail.

Fumaric and maleic acids have been found to condense with p-cresol in the presence of sulfuric acid to give 6-methylcoumarin in good yield. 67.68 The encouraging results in this condensation justify a more

⁶² Rangaswami and Seshadri, Proc. Indian Acad. Sci., 6A, 112 (1937) [C. A., 32, 559 (1938)].

⁶³ Sen and Chakravarti, J. Indian Chem. Soc., 6, 793 (1929).

⁶⁴ Rangaswami and Seshadri, Proc. Indian Acad. Sci., 9A, 7 (1939) [C. A., 33, 4244 (1939)].

⁶⁵ Mentzer, Billet, Molho, and Dat Xuong, Bull. soc. chim. France, 12, 161 (1945) [C. A., 40, 865 (1946)].

⁶⁵ Mentzer and Billet, Bull. soc. chim. France, 12, 292 (1945) [C. A., 40, 2828 (1946)].

⁶⁷ Pondorff, Ger. pat. 338,737 (1921) [C. A., 16, 3488 (1922)].

⁶⁸ Thompson and Edee, J. Am. Chem. Soc., 47, 2556 (1925)

detailed investigation of the condensation of these acids with other phenols.

$$_{\mathrm{H_3C}}$$
 OH + $_{\mathrm{CH-CO_2H}}^{\mathrm{CH-CO_2H}}$ \longrightarrow $_{\mathrm{H_3C}}$ CH

Reactivity of β -Ketonic Esters. Ethyl acctoacetate probably condenses in its enol form with the phenols. β -Ketonic esters with substituents likely to increase the enolization or stabilize the enolic form should therefore be more active than ethyl acetoacetate, and those with substituents that tend to decrease the enolization or lead to a less stable enol form should be less reactive. Substituents in a β -ketonic ester may be attached to the α -carbon atom or the γ -carbon atom, and they provide a means of obtaining coumarins with different substituents in the heterocyclic ring. Cyclic β -ketonic esters, and β -ketonic esters with heterocyclic rings, have also been condensed with phenols. The reactivities of these esters differ very widely.

Ethyl α -chloroacetoacetate has been condensed with a number of phenols to yield 3-chlorocoumarins. The condensation with this ester is smooth and the reactions closely parallel those with ethyl acetoacetate. The corresponding α -bromo ester has not been studied.

In ethyl α -alkyl- and α -aryl-acetoacetates the reactivity varies with the nature of the α substituent. With methyl, ethyl, propyl, butyl, allyl, phenyl, and benzyl groups as α substituents the condensation with reactive phenols is satisfactory, but with less reactive phenols the yields are generally poor and the condensation may be inhibited completely. Thus with m-cresol the α -ethyl derivative of ethyl acetoacetate gives a poorer yield than the α -methyl derivative; α -propyl- and α -phenyl-acetoacetates do not react. The ethyl α -allylacetoacetate, however, condenses with m-cresol easily. He has a home phenols does not react with ethyl α -ethyl-, α -propyl-, or α -isopropyl-acetoacetate. Ethyl α -(α -hydroxy- β , β , β -trichloroethyl) acetoacetate with various phenols gives satisfactory results. Thus the presence of a heavy α substituent like —CH(OH)CCl₃ does not appreciably inhibit the Pechmann reaction and has less effect than an α -ethyl substituent.

The Pechmann reaction of diethyl acetosuccinate and diethyl aceto-

⁶⁹ Chakravarti and Banerjee, J. Indian Chem. Soc., 13, 619 (1936).

Naik, Desai, and Desai, J. Indian Chem. Soc., 6, 83 (1929).
 Chakravarti, J. Indian Chem. Soc., 9, 389 (1932).

Kulkarni, Alimchandani, and Shah, J. Indian Chem. Soc., 18, 113 (1941).
 Kulkarni, Alimchandani, and Shah, J. Indian Chem. Soc., 18, 123 (1941).

⁷⁴ Shah and Kulkarni, J. Univ. Bombay, 10 (pt. 3), 86 (1941) [C. A., 36, 3796 (1942)]

glutarate, which have —CH2CO2C2H5 and —CH2CH2CO2C2H5 as substituents in the α position, with various phenols has been systematically studied. Diethyl acetosuccinate condenses with very reactive phenols and also with m-cresol, 2-acetyl, 2-benzoyl-, and 4-chloro-resorcinol, and 4-chloro-α-naphthol, but not with phenol, o-cresol, p-cresol, hydroquinone, catechol, 4-chlorophenol, \beta-resorcylic acid, resacetophenone. or gallic acid. 34, 42, 75, 76 The presence of a carbethoxyalkyl group as a substituent in the β -ketonic ester results in a molecule of greater reactivity than one in which an alkyl substituent is present; diethyl acetosuccinate is as reactive as or even more reactive than the corresponding ethyl α-alkylacetoacetates. Similar observations have been made with diethyl α-acetoglutarate.77 With substituents such as cyano or aceto the condensation takes place with the elimination of the group and the formation of the unsubstituted coumarin. 32, 46, 78

Other a-substituted ethyl acetoacetates that have been studied are ethyl o-carboxybenzylacetoacetate, 79 ethyl phthalylacetoacetate. 79 ethyl benzoylacetoacetate, 32,46 diethyl acetylmalonate, 32 and ethyl diacetylacetate.32 The first two have been condensed with resorcinol and a few other reactive phenols in the presence of dry hydrogen chloride in acetic acid to form coumarin derivatives. When ethyl benzoylacetoacetate and ethyl diacetylacetate react with resorcinol, the acetyl group is removed during condensation and the same coumarins result as are formed with ethyl benzoylacetate and ethyl acetoacetate, respectively. Diethyl acetylmalonate reacts with the loss of a carbethoxyl group to give the same coumarin as that obtained by the use of ethyl acetoacetate.

A number of β -ketonic esters with groups other than methyl in the γ position have been condensed with phenols. Ethyl butyroacetate. 35 which may be considered as ethyl γ-ethylacetoacetate and ethyl γ-phenylacetoacetate 80,81 react with resorcinol, orcinol, pyrogallol, phloroglucinol, and α-naphthol to give 4-ethyl- and 4-benzyl-coumarin derivatives, respectively, but they do not condense with phenol, β-naphthol, hydroquinone, m-cresol, methyl β -resorcylate, or resacetophenone. A γ substituent thus reduces the reactivity.

Acetonedicarboxylic acid and its diethyl ester have been condensed with a number of simple and substituted phenols.26,46,82 Citric acid gives

⁷⁵ Banerjee, J. Indian Chem. Soc., 8, 777 (1931).

⁷⁶ Dey and Sankarnarayan, J. Indian Chem. Soc., 8, 817 (1931).

⁷ Shah and Shah, Ber., 71, 2075 (1938).

⁷⁸ Held, Compt. rend., 116, 720 (1893).

⁷⁹ Bülow, Ber., 38, 474 (1905).

⁸⁰ Sonn and Litten, Ber., 66, 1512 (1933).

⁸¹ Kotwani, Sethna, and Advani, J. Unir. Bombay, 10 (pt. 5), 143 (1942) [C. A., 37, 623 (1943)].

²² Burton and Pechmann, Ann., 261, 166 (1891).

acetonedicarboxylic acid on heating with concentrated sulfuric acid, and several workers have therefore preferred to condense citric acid with phenols instead of using pure acetonedicarboxylic acid. Phenol, nitrophenols, phenol carboxylic acids, and o- and p-aminophenol have been found not to react. Catechol, o- and p-cresol, hydroquinone, β -naphthol, and methyl β -resorcylate gave poor yields of the corresponding coumarin, but m-cresol, pyrogallol, resorcinol, phloroglucinol, and α -naphthol gave good yields. Thus a molecule with the carboxyl or carbethoxy group in the γ position of ethyl acetoacetate is more reactive than one with a γ -ethyl or γ -phenyl substituent.

Ethyl γ -bromoacetoacetate and m-cresol, α -naphthol, or β -naphthol yield 4-bromomethylcoumarins.⁸³

Among other β -ketonic esters which have been condensed with phenols are ethyl benzoylacetate, 2,13,32,55,65,84 ethyl veratroylacetate, 85,86 diethyl benzoylacetate, 87 diethyl veratroylacetate, 87 diethyl oxalacetate, 26,88,89 diethyl oxalochloroacetate, 26,89 diethyl oxalobromoacetate, and ethyl α -formylphenylacetate. With the exception of diethyl oxalacetate no systematic study has been made with these esters, and no generalizations are therefore possible. Unlike other β -ketonic esters, diethyl oxalacetate either does not condense or gives poor yields with certain meta-substituted phenols but does react more satisfactorily with certain para-substituted phenols; resorcinol and m-cresol give poor yields of coumarins, and orcinol and pyrogallol give no products. Hydroquinone, however, yields the ester of coumarin-4-carboxylic acid.

Several cyclic β-ketonic esters like ethyl cyclopentanone-2-carboxylate ^{36,48,91} and its 4-methyl homolog, ^{48,91,92} ethyl cyclohexanone 2-carboxylate ^{9,48,93,94,95} and its 4-,^{10,36,93,96,97} 5-,^{9,10,38,93,96,97} and 6-^{93,97} methyl homologs, ethyl 3,5-dimethyl-,⁹⁸ ethyl 4,5-dimethyl-,⁹⁸ and ethyl 5,5-dimethyl-cyclohexanone-2-carboxylate,⁹⁸ ethyl cycloheptanone-2-carboxylate,⁹⁸ and ethyl trans-β-decalone-3-carboxylate ^{96,97} have

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El Dey and Sankarnarayan, J. Indian Chem. Soc., 11, 687 (1934).
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⁸⁴ Robinson and Turner, J. Chem. Soc., 113, 874 (1918).

Appel, Baker, Hagenbach, and Robinson, J. Chem. Soc., 1937, 738.
 Mitter and Paul, J. Indian Chem. Soc., 8, 271 (1931).

⁸⁷ Robinson and Rose, J. Chem. Soc., 1933, 1469.

⁸³ Pechmann and Graeger, Ber., 34, 378 (1901).

⁸⁹ Biginelli, Gazz. chim. ital., 24, 491 (1894).

⁹⁰ Huntress and Oleson, J. Am. Chem. Soc., 70, 2831 (1948).

Ahmad and Desai, Proc. Indian Acad. Sci., 5A, 277 (1937) [C. A., 31, 5785 (1937)].
 Dieckmann, Ann., 317, 27 (1901).

Adams, Smith, and Loewe, J. Am. Chem. Soc., 63, 1973 (1941).

Sen and Basu, J. Indian Chem. Soc., 5, 467 (1928).
 Adams and Mecorney, J. Am. Chem. Soc., 66, 802 (1944).

⁸⁶ Chowdhry and Desai, Proc. Indian Acad. Sci., 8A, 1 (1938) [C. A., 32, 9065 (1938)].

⁷ Chowdhry and Desai, Proc. Indian Acad. Sci., 8A, 12 (1938) [C. A., 32, 9006 (1938)].

⁸ Adams, Loewe, Theobald, and Smith, J. Am. Chem. Soc., 64, 2653 (1942).

been condensed with phenols in the presence of sulfuric acid or phosphorus oxychloride. Chowdhry and Desai 97 report that the cyclic β -ketonic esters are more reactive than their open-chain analogs. The sluggishness of ethyl 6-methylcyclohexanone-2-carboxylate as compared with its 4-methyl and 5-methyl analogs may be attributed to the steric hindrance offered by the methyl group in the *ortho* position to the enolic hydroxyl.

Heterocyclic β -ketonic esters like ethyl chroman-3-one-4-carboxylate, 99 ethyl 8-methoxy-, 99 ethyl 3-hydroxy-6,7-dimethoxy-, 99 and ethyl 3-hydroxy-7-methoxy- Δ^3 -chromene-4-carboxylate, 99 ethyl β -coumaranone-2-carboxylate, 100 ethyl 5-methyl-, 100 7-methyl-, 100 and 6-methoxy- β -coumaranone-2-carboxylate, 100 and methyl 3-hydroxyindole-2-carboxylate 100 condense with reactive phenols like resorcinol, phloroglucinol, pyrogallol, and 2-isoamylresorcinol in the presence of sulfuric acid and hydrogen chloride with formation of chromeno- and coumarono-coumarins.

Condensing Agents. The role of the condensing agent in the Pechmann reaction is very important. Condensation between a phenol and a β -ketonic ester that is not brought about in the presence of one condensing agent may be brought about by the presence of another. The yields of product with different reagents may vary markedly. Occasionally one reagent will effect the formation of one type of product and a different reagent an entirely different product.

Of the several condensing agents tested in place of sulfuric acid, only phosphorus pentoxide, phosphorus oxychloride, aluminum chloride, and to some extent zinc chloride have yielded results that require discussion.

Sulfuric Acid and Phosphorus Pentoxide. Simonis 3,4 condensed β -ketonic esters with phenols in the presence of phosphorus pentoxide and reported the formation of chromones exclusively. This conclusion was later found to be incorrect since the condensation product of resorcinol and ethyl α -methylacetoacetate, to which was assigned the structure 7-hydroxy-2,3-dimethylchromone by Simonis and Remmert, 5 was proved by Robertson and his co-workers 101 to be 7-hydroxy-3,4-dimethylcoumarin.

Jacobson and Ghosh condensed various phenols with ethyl α -phenyland α -benzyl-acetoacetate and with ethyl α -benzylbenzoylacetate in the presence of sulfuric acid 102, 103, 104 and reported the products as chromones.

⁹⁹ Hilton, O'Donell, Reed, Robertson, and Rusby, J. Chem. Soc., 1936, 423.

¹⁰⁰ King, Holland, Reed, and Robertson, J. Chem. Soc., 1948, 1673.

¹⁰¹ Canter, Curd, and Robertson, J. Chem. Soc., 1931, 1255.

¹⁰² Jacobson and Ghosh, J. Chem. Soc., 107, 424 (1915).

¹⁰³ Jacobson and Ghosh, J. Chem. Soc., 107, 959 (1915).

¹⁰⁴ Jacobson and Ghosh, J. Chem. Soc., 107, 1051 (1915).

This was due to erroneous interpretation of the results of hydrolysis of the condensation products. Baker 105,100 proved that in the reactions described by Jacobson and Ghosh only coumarins resulted.

An extensive study of the two condensing agents sulfuric acid and phosphorus pentoxide has been made, especially by Robertson 13,107,103 and Chakravarti 33,100 and their eo-workers. From the results obtained so far the following generalizations can be made.

- 1. When sulfurie acid is used as a condensing agent a coumarin is almost always formed. However, β -naphthol and ethyl acctoacetate in the presence of sulfuric acid yield a mixture of a commarin and a chromone in which the eoumarin preponderates.118 From 4-chloro-3,5-dimethylphenol and ethyl acetoacetate a chromone is formed exclusively.95
 - 2. Phenols like resorcinol, pyrogallol, phloroglucinol, orcinol, and α-naphthol that react readily in the presence of sulfurie acid also give coumarins when phosphorus pentoxide is used as the condensing agent.
 - 3. Phenols that do not form coumarins at all or form them in poor yields with sulfurie acid generally give chromones in the presence of phosphorus pentoxide. Thus phenol, o-cresol, halogenated 111 and nitro phenols,29 halogenated and nitro cresols,60 p-xylenol,112 and β -naplithol,110 which either do not condense in the presence of sulfurie acid or eondense with difficulty, are found to give chromones in the presence of phosphorus pentoxide. Some phenols like entechol, for example, do not eondense in the presence of either sulfurie acid or phosphorus pentoxide.
 - 4. With phosphorus pentoxide, chromone formation is favored from β-ketonic esters with an α-alkyl substituent. If the substituent is large, the condensation may be retarded or completely inhibited. m-Cresol and p-cresol with ethyl acetoacetate in the presence of phosphorus pentoxide give the coumarins, 13, 113 but with ethyl α -methyl- and α -ethylacetoacetate they give chromones.3.13,113 Similar results are obtained with 4-chloro- and 4-bromo-α-naphthol.61

Phosphorus Oxychloride. When Naik, Desai, and Desai 70 found that α-naphthol did not condense with ethyl α-benzylacetoacetate in the presence of sulfuric acid they tried phosphorus oxychloride as condensing agent and succeeded in bringing about a reaction. Since then phosphorus oxychloride has been used frequently and in certain cases

¹⁰⁵ Baker, J. Chem. Soc., 127, 2349 (1925).

¹⁰⁶ Baker and Robinson, J. Chem. Soc., 127, 1981 (1925).

¹⁰⁷ Canter, Martin, and Robertson, J. Chem. Soc., 1931, 1877. 108 Robertson, Sandrock, and Hendry, J. Chem. Soc., 1931, 2426.

¹⁰⁹ Chakravarti, J. Indian Chem. Soc., 8, 129 (1931).

¹¹⁰ Dey and Lakshminarayan, J. Indian Chem. Soc., 9, 149 (1932).

¹¹¹ Simonis and Schumann, Ber., 50, 1142 (1917).

¹¹² Goodall and Robertson, J. Chem. Soc., 1936, 426.

¹¹³ Robertson and Sandrock, J. Chem. Soc., 1932, 1180.

successfully where sulfuric acid has failed. 4-Acylresorcinols and gallacetophenone do not condense with ethyl acetoacetate in the presence of sulfuric acid but condense readily in the presence of phosphorus oxychloride to give 6-acylcoumarins. Ethyl 6-methylcyclohexanone-2-carboxylate fails to react with phenols in the presence of sulfuric acid but condenses in the presence of phosphorus oxychloride to give the expected coumarin derivatives. 97

Phosphorus oxychloride frequently gives better yields than sulfuric acid. The condensations of resorcinol, pyrogallol, orcinol, and α -naphthol with diethyl acetosuccinate,³⁴ the condensations of 4-ethyl- and 4-propyl-resorcinol with ethyl α -(α -hydroxy- β , β , β -trichloroethyl)aceto-acetate,⁷⁴ and the condensation of orcinol with ethyl cyclohexanone-2-carboxylate ¹⁰ may be cited as examples.

Although in general phosphorus oxychloride gives the same products as sulfuric acid, the possibility of chromone formation is not precluded. 2-Hydroxy-p-xylene gives rise to chromones on condensation with ethyl α -alkylacetoacetates and ethyl benzoylacetate in the presence of phosphorus oxychloride. 112 4-Hydroxy-m-xylene with ethyl acetoacetate gives 4,6,8-trimethylcoumarin but with ethyl α -methyl- and α -ethyl-acetoacetate yields 2,3,6,8-tetramethyl- and 2,6,8-trimethyl-3-ethyl-chromone, respectively. 112 These are the only instances known of chromone formation in the presence of phosphorus oxychloride. Phosphorus pentoxide gives chromones in each of these reactions.

Anhydrous Aluminum Chloride. In exploring the use of other condensing agents for the Pechmann reaction, Sethna, Shah, and Shah found that anhydrous aluminum chloride dissolved in dry ether or more generally in dry nitrobenzene not only proved to be an efficient condensing agent but also changed the course of some reactions. If the 4 position in resorcinol is occupied by groups such as carboxyl, carbomethoxyl, acyl, or nitro, the condensation instead of giving the 7-hydroxycoumarins gives either exclusively, or mainly, 5-hydroxycoumarin derivatives. These cannot be prepared or can be prepared only with difficulty by any other procedure.

Resacetophenone and other 4-acylresorcinols that do not condense with β -ketonic esters in the presence of sulfuric acid and that give 7-hydroxy-6-acylcoumarins in the presence of phosphorus oxychloride yield 5-hydroxy-6-acylcoumarins in the presence of anhydrous aluminum chloride. 17, 53, 114, 115 The condensation of resacetophenone with cthyl α -methylacetoacetate, which cannot be effected by phosphorus oxychloride, takes place with ethyl α -methyl- and α -ethyl-acetoacetate in

.

Deliwala and Shah, J. Chem. Soc., 1939, 1250.
 Chudgar and Shah, J. Indian Chem. Soc., 21, 175 (1944).

the presence of aluminum chloride. 116 2-Acetylresorcinol and ethyl acetoacetate give the same coumarin and in better yield than with sulfuric acid.17 o-Hydroxyacetophenone, gallacetophenone, quinacetophenone, and resacetophenone with nitro, carbomethoxyl, or aceto substituents, however, do not react with ethyl acetoacetate in the presence of aluminum ehloride.17,117

4-Nitroresorcinol with ethyl acetoacetate in the presence of sulfuric acid yields 7-hydroxy-4-methyl-6-nitroeoumarin," but in the presence of anhydrous aluminum ehloride gives 5-hydroxy-1-methyl-6-nitrocoumarin.118

Methyl β-resorcylate, which condenses with ethyl nectoacetate in the presence of sulfuric acid with formation exclusively of 7-hydroxycoumarin,45 eondenses in the presence of aluminum ehloride to give mainly the 5-hydroxyeoumarin ester and a small quantity of the 7hydroxy isomer.53

With simple phenols the same coumarins are obtained as with sulfurie acid. The yields are higher in some cases and lower in others. Phenol is converted to 4-methylcoumarin in 3% yield on condensation with ethyl acetoacetate in the presence of sulfuric acid,24 but the same coumarin is obtained in 40-55% yield in the presence of aluminum chloride.119

In the condensation of methyl β -resorcylate with ethyl acetoacetate in the presence of zinc chloride, 53 in the condensation of β -resorcylic acid with malic acid in the presence of sulfuric acid,120 and in the condensation of resacetophenone with ethyl acetoacetate in the presence of phosphorus oxychloride,12 5-hydroxycoumarin derivatives have also been isolated in very poor yields, the main products being the 7-hydroxycoumarin derivatives.

¹¹⁶ Deliwala and Shah, Proc. Indian Acad. Sci., 17A, 7 (1943) [C. A., 37, 4379 (1943)]. 117 Deliwala and Shah, Proc. Indian Acad. Sci., 13A, 352 (1941) [C. A., 35, 7959 (1941)].

¹¹⁸ Parekh and Shah, J. Indian Chem. Soc., 19, 339 (1942). 119 Woodruff, Org. Syntheses, 24, 69 (1944).

¹²⁰ Kumar, Ram, and Ray, J. Indian Chem. Soc., 23, 365 (1946).

Zinc Chloride. Zinc chloride has been employed to a very limited extent as a condensing agent.^{32, 121, 122} It does not appear to be superior to phosphorus oxychloride. Generally, the same coumarins are obtained as with sulfuric acid. From methyl β -resorcylate and ethyl acetoacetate in the presence of zinc chloride as the condensing agent, the 7-hydroxycoumarin is the main product with a very small quantity of the 5-hydroxycoumarin.⁵³

Hydrogen Chloride. 62, 79, 85, 99, 100, 123 The advantages of hydrogen chloride as a condensing agent lie in the avoidance of sulfonation of aromatic nuclei, prevention of saponification of the β -ketonic ester, improved yields, and purer products. However, when little or no reaction can be effected with sulfuric acid, as in the case of phenol, β-naphthol, and quinol, hydrogen chloride also gives negative results. In the condensation of ethyl a-allylacetoacetate with phenols a molecule of hydrogen chloride adds at the double bond and, instead of 3-allylcoumarins, 3,β-chloropropylcoumarins are obtained. 70,124 A combination of zinc chloride and hydrogen chloride has been used to advantage 125, 126 in some condensations, especially in those where the other condensing agents give indifferent results. Thus ω-chlororesacetophenone, which did not condense with diethyl oxalacetate in the presence of sulfuric acid or phosphorus pentoxide, did condense in the presence of zinc chloride and dry hydrogen chloride to give β-carbethoxy-6-chloroaceto-7-hydroxycoumarin. 126

Other Condensing Agents. Like hydrogen chloride, phosphoric acid ¹²⁷ is also an effective condensing agent and does not give colored products, but it generally fails to promote condensation where sulfuric acid fails. Other condensing agents that have been reported arc sodium ethoxide, ¹²⁷ boric anhydride, ¹²⁷ sodium acetate, ¹²⁷ ferric chloride, ¹²⁸ stannic chloride, ¹²⁸ titanium chloride, ¹²⁸ and thionyl chloride. ¹²⁹ In the few condensations that have been tried with these reagents, most of them with simple phenols, the same coumarins are obtained as with sulfuric acid. The meager data available do not justify any conclusions regarding their efficacy.

¹²¹ Pechmann and Schwarz, Ber., 32, 3699 (1899).

¹²² Pechmann and Schaal, Ber., 32, 3690 (1899).

¹²³ Appel, J. Chem. Soc., 1935, 1031.

¹²⁴ Ahmad and Desai, J. Univ. Bombay, 6 (pt. 2), 89 (1937) [C. A., 32, 4561 (1938)].

¹²⁵ Borsche and Niemann, Ber., 62, 2043 (1929).

¹²⁶ Gaind, Gupta, Ray, and Sareen, J. Indian Chem. Soc., 23, 370 (1946).

¹²⁷ Chakravarti, J. Indian Chem. Soc., 12, 536 (1935).

¹²³ Horii, J. Pharm. Soc. Japan, 59, 201 (1939) [C. A., 33, 4973 (1939)].

Dixit, Kankudti, and Mulay, J. Indian Chem. Soc., 22, 207 (1945).

EXPERIMENTAL CONDITIONS AND PROCEDURES

The experimental conditions depend on the condensing agent used and are discussed under separate headings. The reaction between certain phenols, especially nitrophenols, and the β -ketonic ester may be Initial heating wherever necessary should therefore be violent.130 gradual.

The ethyl α -alkylacetoacetates may contain ethyl acetoacetate as an impurity. They must be carefully purified, since phenols condense very readily with ethyl acetoacetate and a mixture of coumarins may result from which a pure product may be difficult to isolate. Ethyl acetoacetate may be removed from the α -alkyl derivatives by washing with 3% sodium hydroxide solution. The washed product is then distilled.47 This method is more satisfactory than fractional distillation under reduced pressure, especially for ethyl α -methyl- and α -ethyl-acetoacetate contaminated with ethyl acetoacetate.

Sulfuric Acid as Condensing Agent

Concentrated sulfuric acid is generally used as the condensing agent. However, 73-80% sulfuric acid is sometimes preferable as it will decrease the tendency to sulfonation. The addition of the sulfuric acid to the mixture of phenol and \beta-ketonic ester should be gradual, preferably with cooling, since sufficient heat may be evolved to char the product. The reaction mixture is allowed to stand overnight or for a number of days, depending on the reactivities of the phenol and the β -ketonic ester used. After the required period the reaction mixture is added slowly to cold water or crushed ice and the coumarin is precipitated. Sometimes, after the addition of sulfuric acid to the mixture of phenol and β -ketonic cster, the reaction mixture may be heated on a steam bath for some time, and then left at room temperature for one or more days. Reactions are also described in which heating on the steam bath is started immediately and continued for three to four hours, after which the reaction mixture is cooled and added to ice water. Condensations that proceed with difficulty, such as those of phenols with malic acid, are usually carried out at temperatures up to 150°. 6-Methylcoumarin was synthesized best by mixing the cresol and sulfuric acid, maintaining the mixture in a bath at 135°, and introducing the malic acid slowly.¹³¹ The yield is generally low when heating is required, since a portion of the product may be sulfonated.

7-Hydroxycoumarin.8 An intimate mixture of 3 g. of resorcinol, 2.46 g. of malic acid, and 6.1 ml. of concentrated sulfuric acid, after

¹¹⁰ Chakravarti, J. Indian Chem. Soc., 9, 25 (1932).

in Bailey and Boettner, J. Ind. Eng. Chem., 13, 905 (1921).

200

being heated in an oil bath at 120° until the effervescence ceases (one hour), is cooled and treated with excess of crushed ice. The precipitated coumarin is purified by repeated crystallization from dilute ethanol (decolorizing carbon), from which it separates as pale pink prisms, m.p. 227-228°; yield 43%. The crude product can be conveniently decolorized by passing a stream of sulfur dioxide into a warm ethanolic solution

The success of the method, according to Dey, Rao, and Seshadri, 132 depends primarily on the regulation of the heating. It should be stopped precisely at the moment the mixture becomes clear.

7-Hydroxy-4-methylcoumarin. 133 The preparation of this coumarin from resorcinol and ethyl acetoacetate with concentrated sulfuric acid as the condensing agent has been described in Organic Syntheses. The yield is 82-90%.

6,7-Dihydroxy-4-methylcoumarin. 60 The preparation of this coumarin from 1,2,4-triacetoxybenzene and ethyl acetoacetate in the presence of 75% sulfuric acid has been described in Organic Syntheses. The yield is 92%.

Phosphorus Pentoxide as Condensing Agent

The condensation may be carried out in the presence of this agent either in the cold if the phenol is very reactive or by heating the reaction mixture if the phenol is less reactive. The initial reaction is very vigorous, and external cooling is essential. It has been observed that the addition of a little absolute ethanol is advantageous.33

5-Hydroxy-4,7-dimethylcoumarin.33 To a mixture of 5 g. of orcinol and 5 g. of ethyl acetoacetate cooled in ice, 18 g. of phosphorus pentoxide is added gradually. A vigorous reaction takes place with evolution of much heat. When the reaction ceases, the cold mass is treated with water. The precipitate is washed with water and crystallized from dilute ethanol (decolorizing carbon). It forms colorless needles, m.p. 248°.

2,5-Dimethyl-3-ethylchromone. The vigorous reaction between 20 g. of m-cresol, 5 g. of ethyl α -ethylacetoacetate, and 20 g. of phosphorus pentoxide is controlled by agitation and occasional cooling in tap water. Then a further 10 g. of m-cresol and 20 g. of the pentoxide are added. The mixture is heated at 140° in an oil bath for fifteen minutes and then on the steam bath for one hour. An aqueous solution of the dark-colored product is made basic with sodium hydroxide and extracted with ether. After the evaporation of the solvent the extract is distilled under reduced pressure and the main fraction, b.p. 170-190°/20 mm., is mixed with an equal volume of light petroleum ether. 2,5-Di-

in Dey, Rao, and Seshadri, J. Indian Chem. Soc., 11, 746 (1934).

in Russell and Frye, Org. Syntheses, 21, 22 (1941).

mcthyl-3-ethylchromone, which gradually crystallizes, is separated; and, after the removal of the solvent, the mother liquor is distilled in a vacuum. When the distillate is mixed with petroleum ether a further quantity of the solid is obtained. On recrystallization from the same solvent, the chromone forms thick, pointed prisms, m.p. 86°; yield, 1 g.

Phosphorus Oxychloride as Condensing Agent

Dry benzene or toluene is generally the solvent when phosphorus oxychloride is used as condensing agent. The reaction mixture is usually heated for a few hours on a steam bath.

7-Hydroxy-4-methyl-6-acetylcoumarin and 5-Hydroxy-4-methyl-6-acetylcoumarin. A mixture of 8 g. of resacetophenone, 6 g. of ethyl acctoacetate, 2 ml. of phosphorus oxychloride, and 20 ml. of dry benzene protected from moisture is heated on a steam bath for five hours, when the evolution of hydrogen chloride ceases. After the benzene solution is poured off, the residue is extracted with two portions of 20 ml. of benzene and the solvent is removed by distillation from the combined extracts. The residue obtained from the benzene extracts is recrystallized from ethanol, and pure crystals of 7-hydroxy-4-methyl-6-acetylcoumarin, m.p. 212°, are obtained. The yield is 40%. Concentration of the ethanolic mother liquor gives a second crop of lower purity. The residue left after the removal of the solvent is repeatedly extracted with petroleum ether (b.p. 60-80°). Upon cooling, crystals deposit which on recrystallization from ethanol yield 5-hydroxy-4-methyl-6-acetylcoumarin, m.p. 164-165°.

1-Hydroxy-3-methyl-7,8,9,10-tetrahydro-6-dibenzopyrone. A solution of 6.2 g. of orcinol, 11 g. of ethyl cyclohexanone-2-carboxylate, and 4.6 ml. of phosphorus oxychloride in 45 ml. of dry benzene in an all-glass apparatus and protected from moisture is refluxed for three hours on the steam bath. The solution rapidly turns deep red, and at the end of one hour a crystalline precipitate begins to separate. Two volumes of water are added; the mixture is well shaken to destroy the phosphorus oxychloride and then cooled. Most of the product crystallizes and is obtained by filtration of the benzene-water mixture. Additional material is obtained by separation and evaporation of the benzene layer. Purification is effected by recrystallization from ethanol, m.p. 243-245°; yield, 7.6 g. (66%).

Anhydrous Aluminum Chloride as Condensing Agent

Anhydrous aluminum chloride can be used as the condensing agent either without added solvent or dissolved in dry ether or dry nitrobenzene. The best results have been reported with nitrobenzene. The aluminum ehloride is dissolved in dry, preferably freshly distilled nitrobenzene, by warming in a flask protected from moisture. This solution is added to the solution of the phenol and the β -ketonie ester in dry nitrobenzene. The reaction mixture is heated in an oil bath between 120° and 140° for an hour or two, when the evolution of hydrogen ehloride almost ceases. At the end of that period the reaction mixture is eooled and the unused aluminum chloride is decomposed by the addition of ice and eoneentrated hydroehloric acid. The nitrobenzene is removed by steam distillation. The product remains behind. It is generally found that two moles of aluminum chloride per mole of the phenol give the best yield; more or less aluminum ehloride than this quantity may decrease the yield. The pure anhydrous aluminum chloride dissolves in ether and nitrobenzene without leaving a residue.

Methyl 5,7-Dihydroxy-4-methylcoumarin-6(or 8)-carboxylate. Two grams of methyl phloroglueinolearboxylate and 1.5 g. of ethyl acetoacetate are dissolved in a minimum quantity of dry ether. To this solution 3.5 g. of anhydrous aluminum chloride in 15 ml. of dry ether is added. The ether is allowed to evaporate gradually by heating the flask on a warm water bath, and the resulting homogeneous mass is heated in an oil bath between 120° and 125° for an hour until the evolution of hydrogen chloride is negligible. After cooling, dilute hydrochloric acid and ice are added. The product is purified by crystallization from ethanol. It forms clusters of tiny needles, m.p. 230-231°; yield, 1.2 g.

5-Hydroxy-4-methyl-6-propionylcoumarin.¹¹⁴ A solution of 4.2 g, (1 mole) of anhydrous respropiophenone and 3.25 g. (1 mole) of ethyl acetoacetate in dry nitrobenzene is added to a solution of 6.7 g. (2 moles) of anhydrous aluminum ehloride in 35 ml. of dry nitrobenzene. The mixture, protected from moisture, is heated at 120-130° until evolution of hydrogen chloride is negligible, which takes about an hour. It is then cooled, ice and 15 ml. of concentrated hydrochloric acid are added, and the nitrobenzene is steam-distilled. The brown residue is collected, decolorized by washing with a small quantity of ethanol, and crystallized from ethanol. It forms fine, silky needles, m.p. 164-165°; yield, 2 g.

Hydrogen Chloride as Condensing Agent

A solution of the phenol and the β -ketonic ester either in glacial acetic acid or in absolute ethanol is a saturated with hydrogen chloride while being cooled with ice water, and the reaction mixture is kept in a well-stoppered flask overnight. It is then poured into water directly or after heating for some time on a steam bath. The commarin precipitates.

7-Hydroxy-5'-methylcoumarono-(2',3',3,4)-coumarin. When a solution of 1 g. of ethyl 5-methyl-3-coumaranone-2-carboxylate and 1 g. of

resorcinol in methanol is saturated slowly at room temperature with hydrogen chloride a yellow solid gradually separates. After two days the mixture is heated on the steam bath for half an hour, then cooled, and the resulting coumarin is collected, washed, and crystallized from ethanol, m.p. above 300°; yield, 0.6 g.

Zinc Chloride as Condensing Agent

The condensation in the presence of zinc chloride may be carried out either with ethanol as solvent or without a solvent. Heating is essential, the period dependent on the reactivities of the phenol and the β -ketonic ester.

Ethyl 7-Dimethylaminocoumarin-4-acetate.¹³⁴ A mixture of 7 g. of distilled diethyl acetonedicarboxylate, 5 g. of m-dimethylaminophenol, 6 g. of powdered anhydrous zinc chloride, and 20 ml. of absolute ethanol is heated in a paraffin bath with refluxing for twelve hours. The resulting strongly fluorescent liquid, which deposits a small amount of a viscid solid on cooling, is poured into 400 ml. of cold water containing a little hydrochloric acid. A dark oil is precipitated, which, after it has been washed with water containing dilute hydrochloric acid and permitted to stand in contact with ethanol, solidifies slowly to a crystalline cake. The solid is crystallized first from a mixture of benzene and petroleum ether and then from absolute ethanol (decolorizing carbon). The product forms slender, colorless prisms, m.p. 133°. The yield is poor.

TABULAR SURVEY OF THE PECHMANN REACTION

All the condensations of malic acid and β -ketonic esters with phenols and miscellaneous compounds which, in the presence of various condensing agents, have resulted in the formation of either coumarins or chromones have been listed. The literature survey is complete to January, 1949.

The condensations with monohydric phenols are listed in Table I, with dihydric phenols in Table II, with trihydric phenols in Table III, with naphthols in Table IV, and with miseellaneous compounds in Table V.

The condensations with phenol itself are followed by those with monosubstituted phenols with the substituents in the following order: halogens, nitro, amino, alkyl groups in the order of increasing complexity, carboxyl and carbomethoxyl, and acyl. Then are listed the condensations with disubstituted phenols with the substituents in the same

¹³⁴ Dey, J. Chem. Soc., 107, 1643 (1915).

TABLE I
Condensations with Monohydric Phenols

		Condensations	HTIW	MON	OHIDINO 1	Tr 13	Refer	
			Condensing	•		Yield	SOUS	
			-	6	Product	%	-	
Phenol		Acid or Ester	Agent			Poor	1, 14	2
Phenol	Malio	acid	H2SO4	Cour	arm			
			(73% &					
			coned.)				142	
	α-M	ethylmalic acid	H2SO4	3-Me	thylcoumarin			
		•	(73%)			Low	143	
	Eth	yl α-methylformylacetate	P2O5		ethylchromone	3	2, 2	4
		yl acetoacetate	H ₂ SO ₄	4-M	ethylcoumarin	21	14	Į.
		yl acetoacetate	H2SO4	4-M	ethylcoumarin			
			(73%))		2		5
	Et	nyl sodioacetoacetate	P205		lethylchromone	40-55	: 11	9
		hyl acetoacetate	AlCla	4-N	[ethylcoumarin	40-04	14	
		hyl a-methylacetoacetate	H2SO4	3.4	Dimethylcoumarin	_	-	· -
		Til a metal metal metal	(73%					3
	F	thyl c-methylacetoacetate	P2O5	2.3	-Dimethylchromone	25		4
		thyl a-ethylacetoacetate	P2O5	2-1	Methyl-3-ethylchromone		. 4	38
		cetonedicarboxylic acid	H ₂ SO.	. Cr	umarin-4-acetic acid	12		do.
		cetonecocar pozytie acta	1,200	2-	Hydroxyphenylgiutaconic anhydri	de —		45
		Titric acid	H ₂ SO	, c	oumarin-1-acetic acid	7	, ,	24
		Diethyl oxalacetate	H ₂ SO	. E	thyl coumarin-4-carboxylate		•	
		Diethyl ozalochloroacetate	_	. F	thyl 3-chlorocoumarin-4-carboxyla	ite 1		90
		Diethyl oxslobromoacetate). F	thyl 3-bromocoumarin-4-carboxyl	ate 1		90
		Ethyl cyclopentanone-	H ₂ S0). i	Cyclopenteno-(1',2',4,3)-coumarin		5	91
		2-carboxylate	11200	,	3,4ctopenseno-(x 12 1210) 00 200-			
		Ethyl cyclopentanone-	P20		Cyclopenteno-(1',2',2,3)-chromone	-	-	11
		2-carboxylate	1 20	•	Cyclopenteno-(1,2,2,0)-cmcom			
e-Chlo		Ethyl a-methylacetoscet	ste P20	١.	8-Chloro-2,3-dimethylchromone	:	27	111
	nol	Ethyl a-ethylacetoscetat	_		8-Chloro-2-methyl-3-ethylchromon	ie -	-	111
ршe	111/4	Ethyl a-propylacetosceta	-	•	8-Chloro-2-methyl-3-propylchrom		30	130
		Ethyl a-isopropylacetoac			8-Chloro-2-methyl-3-isopropylchro	mone .		130
o-Bro	2000	Ethyl a-methylacetoscei			8-Bromo-2,3-dimethylchromono		17 1	11,130
	nenol	Ethyl a-ethylacetosceta		0 ₅	8-Bromo-2-methyl-3-ethylchromo	ne	23	111
P.	1000	Ethyl a-propylacetoace		O ₆	8-Bromo-2-methyl-3-propylchrom			130
	hloro-	Malic acid		2SO4	7-Chlorocoumarin	10110	4	25
	benol	Ethyl acetoscetate		250t 250t	7-Chloro-4-methylcoumarin		6	25
P-	DCTO!	Ethyl a-methylacetoso		200£	7-Chloro-2,3-dimethylchromone		23	111
		Ethyl a-ethylacetoscet		20s 20s	5 (or 7)-Chloro-2-methyl-3-ethyl-	_	20	111
		zmjez emjacewace	200 I	205	chromone	-		
F7-	Bromo-	Ethyl a-methylacetoa	cetate T	205	5-Bromo-2,3-dimethylchromoos		22	111
	phenol				(7-bromo isomer also formed	but not		
	F				(betalous isomet and formed	DIL MOT		
		Ethyl a-ethylacetoso	tate	P2O5	5 (or 7)-Bromo-2-methyl-3-ethy	chenmona	20	111
D-	Chloro-	Malic acid		H.50.	6-Chlorocoumarin	itmomono.	3	25
•	phenol	Ethyl acetoscetate		H-SO.	6-Chloro-4 methylcoumarin		3	25
	•	Ethyl a-methylaceto		P205	6-Chloro-2,3-dimethylchromon		17	111, 130
		Ethyl a ethylacetose		P2O5	6-Chloro-2-methyl-3-ethylchron			111
		Ethyl a-propylaceto		P2O3	6-Chloro-2-methyl-3-propylchr			130
		Ethyl a-isopropylac			6-Chloro-2-methyl-3-isopropyle			130
		Diethyl acetonedica	uboxylate	H-SO			<6	26
		Diethyl omlacetate		H2504			_	26
		Ethyl cyclopentano	ne-	P203	6-Chloro-2,3-dihydropentachro		Poor	146
		2-carboxylate					_	
	p-Bromo			H;80;			_	144
	Lpeso			(737	76)			
		Ethyl a-methylao		P2O1	6-Bromo-2,3-dimethylchromo	ne	_	111
		Ethyl a-ethylacti	orași e	P2O3	6-Bromo-2-methyl-3-ethylchi	omone	16	111
	11-4-4	D. fanan me 147-244 ava U-		*7 *0	•			

Note: References 142-244 are listed on pp. 57-58.

TABLE I-Continued

CONDENSATIONS WITH MONOHYDRIC PHENOLS

		Condensin	g	Yie	ld Refer-
Pheaol	Acid or Ester	Agent	Product	%	
m-Nitro-	Ethyl a-methylacetoacetate	P2Os	7-Nitro-2,3-dimethylchromone	_	
pheaol	Ethyl α-ethylacetoacetate	P_2O_5	7-Nitro-2-methyl-3-ethylchromone	_	29
	Ethyl α-propylacetoacetate	P_2O_5	7-Nitro-2-methyl-3-propylchromone		
	Et hyl α-isopropylacetoacetate	P ₂ O ₅	7-Nitro-2-methyl-3-isopropylchromone		29
•••	Ethyl a-isohutylacetoacetate	P_2O_5	7-Nitro-2-methyl-3-isohutylchromone	_	29
p-Nitrophenol	•	P ₂ O ₅	6-Nitro-2,3-dimethylchromone	_	29
	Ethyl α-ethylacetoacetate	P ₂ O ₅	6-Nitro-2-methyl-3-ethylchromone		29
	Ethyl a-propylacetoacctate	P_2O_5	6-Nitro-2-methyl-3-propylchromone	-	29
m-Amino-	Ethyl α-isobutylacetoacetate Ethyl acctoacetate	P_2O_5 $ZnCl_2$	6-Nitro-2-methyl-3-isohutylchromone	- 10.1	29
pheaol	Ethyl acctoacetate	ZIICI2	7-Amino-4-methylcoumaria with vary- ing proportions of 7(?)-hydroxy-	12-1	6 121
Phonor			lepidone, 7(?)-hydroxy-2,4,4-tri-		
			methyl-3,4-dihydroquinoline, and		
			4,6,6,8-tetramethyl-6,7-dihydro-		
			quiaocoumarin		
m-Methyl-	Ethyl acetoacetate	$ZnCl_2$	7-Methylamiao-4-methylcoumaria	65	147
amino-					
pheaol	7707 1	. A1			
m-Dimethyl-	Ethyl acetoacetate	ZaCl ₂	7-Dimethylamino-4-methylcoumaria	70-75	
amiao- phenol	Ethyl a-ethylacetoacetate	ZnCl ₂	7-Dimethylamino-3-ethyl-4-methyl- coumaria	_	122
	Diethyl acetonedicarhoxylate	$ZaCl_2$	Ethyl 7-dimethylamiaocoumarin-		26
			4-acetate		
m-Diethyl	Ethyl acetoacetate	$ZnCl_2$	7-Diethylamiao-4-methylcoumaria		122
amino-					
phenol o-Cresol	Ethyl acetoacetate	P ₂ O ₅	2,8-Dimethylchromone	۰	
0-010301	Ethyl a-methylacetoacetate	P ₂ O ₅	2,3,8-Trimethylchromone	8 40	4
	Ethyl a-ethylacetoacetate	P ₂ O ₆	2,8-Dimethyl-3-ethylchromone	-40	130
	Acetonedicarhoxylic acid	H ₂ SO ₄	8-Methylcoumarin-4-acetic acid	25	138
			β-2-Hydroxy-3-methylphenylglutaconic anbydride	_	
	Diethyl acetonedicarhoxylate	H ₂ SO ₄	Ethyl 8-methylcoumarin-4-acetate	_	26
m-Cresol	Malic acid	H ₂ SO ₄	7-Methylcoumaria	27-40	27, 148
	Malie acid	H ₂ SO ₄	7-Methylcoumarin	54	131
		(96%)			
	Ethyl acetoacetate	H_2SO_4	4,7-Dimethylcoumaria *	71	27
	Ethyl acetoacetate	P ₂ O ₅	4.7-Dimethylcoumaria	8	13
	Ethyl a-chloroacetoacetate	H ₂ SO ₄	3-Chloro-4,7-dimethylcoumarin	40	26
	Ethyl α-methylacetoacetate Ethyl α-methylacetoacetate	H ₂ SO ₄ P ₂ O ₅	3.4.7-Trimethylcoumarin 2.3.7-Trimethylchromone	10	27 3
	Ethyl \alpha-methylacetoacetate	P ₂ O ₅	2,3,5-Trimethylchromone	4	13
			2,3.7-Trimethylchromone (isolated as		
			the styryl derivative)		
	Ethyl α-ethylacetoacetate	H ₂ SO ₄	3-Ethyl-4,7-dimethylcoumarin	-	28
	Ethyl α-ethylacetoacetate	P_2O_5	2,5-Dimethyl-3-ethylchromone	2	13
			2,7-Dimethyl-3-ethylchromone (isolated as the styryl derivative)		
	Ethyl α-allylacetoacetate	H ₂ SO ₄	3-Allyl-4,7-dimethylcoumarin	54	70
		H ₂ SO ₄	3-Benzyl-4,7-dimethylcoumarin	-	70 28, 105
		H ₂ SO ₄	Ethyl 4,7-dimethylcoumarin-3-acetate	25	34, 65
		-	4,7-Dimethylcoumarin-3-propionic acid	20	77
		(78%)			

Note: References 142-244 are listed on pp. 57-58.

If the quantity of sulfuric acid employed is less than that given in ref. 27, 4-tolyloxy-4,7-dimethylhydrocoumarin is obtained along with 4,7-dimethylcoumarin, ref. 28.

TABLE I-Continued

CONDENSATIONS WITH MONOHYDRIC PHENOLS

	Condensations	WITI	H INTON	OHYDRIC PHENOUS			
		Condensi	ino		Yield	Ref	
206 1	Acid or Ester	Agent	_	Product	%	en	
Phenol		H ₂ SO ₄		thyl-4-bromomethylcoumarin	_	-	3
rs-Cresol	Ethyl 7-bromoacetoacetate	H ₂ SO ₄	7.35	ethylcoumarin-4-acetic acid	_	_	8
(Confd)	Acetonedicarboxylic acid	H ₂ SO ₄		thylcoumarin-4-acetic acid	60	13	38
	Acetonedicarboxylic acid	проц	β-2-	Hydroxy-4-methylphenylglutaconi hydride	c —		
	Diethyl acetonedicarboxylate	H ₂ SO ₄	7-M	ethylcoumarin-4-acetic acid and i thyl and m-tolyl esters	ta 32-43		49
	Citric scid	H ₂ SO,	7-M	lethylcoumarin-4-acetic acid	44 24	1	.50
	m	77 50	-	Dimethylcoumarin Dimethylcoumarin	8	1	51
	Citric acid (hydrated)	H ₂ SO H ₂ SO		Dimethylcoumarin	4	1	151
	Citric acid (dehydrated)	Oleun		-Dimethylcoumarin	1		151
	Citric acid			-Limethylcoumarin Phenyl-7-methylcoumarin	_		13
	Ethyl benroylscetate	H ₂ SO		Methylffavone	2		13
	Ethyl benroylacetate	P ₂ O ₅			te Poor		26
	Diethyl oxalacetate	H2SC		hyl 7-methylcoumarin-4-carboxyla			26
	Diethyl chloroöxalacetate	H ₂ S0	J4 E	thyl 3-chloro-7-methylcoumarin- 4-carboxylate			
	Ethyl cyclopentanone-	H ₂ S	04 7-	Methylcyclopenteno-	9		91
	2-carboxylate		•••	(1',2',4,3)-coumarin			
	Ethyl cyclopentanone-	P20	s 7.	Methylcyclopenteno-	_		11
	2-carboxylate	- • -	• .	(1',2',2,3)-chromone			
	Ethyl cycloheranone- 2-carboxylate	H ₂ S	SO ₄ 3	.4-Tetrahydrobenzo-7-methylcoum	arin 50	, ()4, 152
r-Tolyl	Ethyl acetoscetate	H-9	304 4	,7-Dimethylcoumarin	_		13
methyl	2103.0001000		86%)	,,-Dimeny.			
ether		•	50707				
p-Cresol	Fumarie acid		SO ₄ ;	8-Methylcoumarin	5	0	67
	Fumarie acid	H		6-Methylcoumarin	40-	80	68, 153
	Maleic acid	H	2504; ZaCl2	6-Methylcoumarin	5	60	67
	Malic acid		SO:	6-Methylcoumarin	•	32	148
	Ethyl acetoscetate		250t	4,5-Dimethylcoumarin		40	2, 28,
			2004	4,0-Dimerny isodinarin	=		154
	Ethyl acetoscetate	F	(80%)	4,6-Dimethylcoumarin		70	155
	Ethyl acetoacetate	ī	201	4.6-Dimethylcoumarin		_	113
	Ethyl acetoscetate		H ₂ PO ₄	4,6-Dimethylcoumsrin		_	127
	Ethyl o-chloroscetoscet		H ₂ SO.	3-Chloro-4,6-dimethylcoumarin		_	26
	Ethyl a-chloroscetoscet		P ₂ O ₄	3-Chloro-4,6-dimethylcoumsrin		_	13
	Ethyl a-methylacetosce		H ₂ SO ₄	3,4,6-Trimethylcoumarin		72	130
	Ethyl o-methylacetoso	tate	(80%)	3,4,6-Trimethylcoumarin			103
	Fibyl o-methylacetoso		P=0;	2.3,6-Trimethylchromone		20	3
	Ethyl a-ethylacetoacet		(\$455)	3-Ethyl-4,6-dimethylcoumarin		7	113
	Ethyl a-ethylacetosce	se	P2Os	2,6-Dimethyl-3-ethylchromone		_	113, 15
	Ethyl o-(a-bydroxy-f.	sstir	H.SO.	4.5-Dimethyl-3-(a-hydroxy-8.8.	8-tri-	18	73
	thlorosthyf)acetoace	tate	•	chloroethyl)coumarin	- ···		
	Dethyl o-acrtylglatar		H:50,	4.6-Dimethylcoumarin-3-propios	nie scid	14	77
	Arricard carboxylic s	-: 4	H2SO.	6-Methyloumarin-4-acetic acid	1	20	26
	Acritocileurocyte s	ಪತ	H-SO.	6-Methylcoumarin-4-acetic acid		40	138
	_			6-2-Hydroxy-5-methylphenylgi- anhydride		_	

TABLE I—Continued Condensations with Monohydric Phenols

7 0.		Condensit	-	Yiel	
Phenol	Acid or Ester	Agent	Product	%	ence
p-Cresol	Citric acid	H ₂ SO ₄	4.6-Dimethylcoumarin	1	151
(Cont'd)	Ethyl benzoylacetate	H ₂ SO ₄ (84%)	4-Phenyl-6-methylcoumarin	2	113
	Ethyl benroylacetate	P2O5	6-Methyldavone	_	113
	Diethyl oxalacetate	H ₂ SO ₄	Ethyl 6-methylcoumarin-4-carboxylate		26
	Ethyl cyclopentanone- 2-carboxylate	H ₂ SO ₄	6-Methylcyclopenteno- (1',2',4,3)-cournarin	8	91
	Ethyl cyclopentanone- 2-carboxylate	P ₂ O ₅	6-Methylcyclopenteno- (1',2',2,3)-chromone	_	11
3-n-Amyl- phenol	Ethyl cyclohexanone- 2-carboxylate	H ₂ SO ₄	3-n-Amyl-7,8,9,10-tetrahydro-6-di- benzopyrono	28	157
	Ethyl 5-methylcyclohexa- none-2-carboxylate	H ₂ SO ₄	3-n-Amyl-9-methyl-7,8,9,10-tetrahydro 6-dibenzopyrone	- 32	157
m-Hexyl- phenol	Malic acid	H ₂ SO ₄	7-Hexylcoumarin	39	158
2,4-Dichloro-	Ethyl α-methylacetoacetate	P_2O_5	6,8-Dichloro-2,3-dimethylchromone	15	111
phenol	Ethyl a-ethylacetoacetate	P2O5	6,8-Dichloro-2-methyl-3-ethylchromone		111, 130
2,4-Dihromo- phenol	Ethyl &-methylacetoscetate	P ₂ O ₅	6,8-Dibromo-2,3-dimethylchromone	19	111
2-Chloro-	Ethyl acetoacetate	H ₂ SO ₄	8-Chloro-4,6-dimethylcoumarin	_	69
4-methyl-	Ethyl a-chloroacetoacetate	H ₂ SO ₄	3,8-Dichloro-4,6-dimethylcoumarin	_	69
phenol	Ethyl a-methylacetoacetate	H ₂ SO ₄	8-Chloro-3,4,6-trimethylcoumarin	_	69
	Ethyl a-methylacetoacetate	P2O5	8-Chloro-2,3,6-trimethylchromone	_	69
	Ethyl α-ethylacetoacetate	H ₂ SO ₄	8-Chloro-3-ethyl-4,6-dimethylcoumarin	_	69
4.00	Ethyl α-ethylacetoacetate	P2O5	8-Chloro-2,6-dimethyl-3-ethylchromone	_	69
4-Chloro-	Ethyl acetoacetate	P ₂ O ₅	6-Chloro-2,8-dimethylchromone	_	69
2-methyl-	Ethyl a-methylacetoacetate	P ₂ O ₅	6-Chloro-2,3,8-trimethylchromone	_	69
phenol	Ethyl a-ethylacetoacetate	P ₂ O ₅	6-Chloro-2,8-dimethyl-3-ethylchromone	_	69 69
	Ethyl α-propylacetoacetate	P ₂ O ₅	6-Chloro-2,8-dimethyl-3-propylchro- mone	_	
4-Chloro-	Ethyl acetoacetate	H ₂ SO ₄	6-Chlore-4,7-dimethylcoumarin	_	69
3-methyl-	Ethyl a-chloroacetoacetate	H ₂ SO ₄	3,6-Dichloro-4,7-dimethylcoumarin	17	69, 159
phenol	Ethyl a-methylacetoacetate	H ₂ SO ₄	6-Chloro-3,4,7-trimethylcoumarin	_	69
	Ethyl a-methylacetoacetate	P2O5	6-Chloro-2,3,7-trimethylchromone	_	69
	Ethyl a-ethylacetoacetate	H ₂ SO ₄	6-Chloro-3-ethyl-4,7-dimethylcoumarin	_	69
	Ethyl a-ethylacetoacetate	P ₂ O ₅	6-Chloro-2,7-dimethyl-3-ethylchromone	_	69
	Ethyl a-propylacetoacetate	P ₂ O ₅	6-Chloro-2,7-dimethyl-3-propylchro- mone	_	69
	Diethyl acetosuccinate	H₂SO₄	Ethyl 6-chloro-4,7-dimethylcoumarin- 3-acetate 6-Chloro-7-methylcoumarin-4-acetic	16	69
	Acetonedicarboxylic acid	H ₂ SO ₄	acid Ethyl 6-chloro-7-methylcoumarin-	Excel-	26, 69 26
2-Nitro-	Diethyl oxalacetate	H ₂ SO ₄ P ₂ O ₅	4-carboxylate 8-Nitro-2,7-dimethylchromone	lent	69
	Ethyl acetoacetate	P ₂ O ₅	8-Nitro-2,7-dimethyl-3-ethylchromone	_	69
3-methyl- phenol	Ethyl a-ethylacetoacetate	P ₂ O ₅	6-Nitro-2,3,8-trimethylchromone		69
4-Nitro-	Ethyl a-methylacetoacetate		6-Nitro-2,8-dimethyl-3-ethylchromone	_	69
2-methyl-	Ethyl α-ethylacetoacetate	P ₂ O ₅ P ₂ O ₅	6-Nitro-2,8-dimethyl-3-propylchromone	_	69
phenol	Ethyl a-propylacetoacetate	H ₂ SO ₄	6,7-Dimethylcoumarin	_	25
3,4-Xyleno1 (3,4-di-	Malic acid	H ₂ SO ₄	4,6,7-Trimethylcoumarin	58	25
methyl- phenol)	Ethyl acetoacetate Ethyl α-chloroacetoacetate	H ₂ SO ₄	3-Chloro-4,6,7-trimethylcoumarin	Very good	26
pnenoi)	Ethyl a-methylacetoacetate	H ₂ SO ₄	3,4,6,7-Tetramethylcoumarin	46	25
	Acetonedicarboxylic acid	-,	6,7-Dimethylcoumarin-4-acetic acid	_	26

Note: References 142-244 are listed on pp. 57-58

ORGANIC REACTIONS

TABLE I-Continued

Condensations with Monohydric Phenols

		CONDENSATIONS	WI'	гн Мо	NOHYDRIC PHENOLS		n (
		C	anđe	nsing		Tierr	Refer- ence	
Phenol		Acid or Ester	Age	nt	Product	%	26	
	D: AL		12SO	. Et	hyl 3-chloro-6,7-dimethylcoumarin-	29	20	
3,4-Xylenol	Dietn	yl chiorooxalaceate	-200		4-carboxylate			
(3,4-di- methyl-								
bpeaol)								
(Cont'd)							160	
2,3-Xylenol	Ethy	l a-methylacetoacetate	P20	ь 2	3.7.8-Tetramethylchromone	_	_	
(2,3-di-								
methyl-								
phenol)					n mt - it it i i i i i i	30	25	
2,4-Xylenol		ie acid	H ₂ S	-	5.8-Dimethylcoumarin	50-97	25, 1	61
(2,4-di-	Eth	yl acetoacetate		SO4 concd.	4.6.8-Trimethylcoumarin			
methyl-				and				
phenol)				36%)				
	T.	hyl acetoacetate		O _b	2,6,8-Trimethylchromone	12-18	16	
		hyl acetoacetate		OCl ₂	4.6.8-Trimethylcoumarin		11	
		hyla-methylacetoacetate		2504	3.4.6.8-Tetramethylcoumarin	25	25,	701
			-	(coned.				
				and				
				86%)			1	61
		thyl a-methylacetoscetate		20ε	2,3,6,8-Tetramethylchromone	16		12
		thyl a-methylacetoscetate		OCI3	2,3,6,8-Tetramethylchromone	_	_	61
	1	Ethyl a-ethylacetoacetate	F	12804	4.6.8-Trimethyl-3-ethylcoumarin	_	-	
		DO 1		(86%)	0.00M1 H 10 H 11		1	161
		Ethyl a-ethylacetoacetate Ethyl a-ethylacetoacetate		P2Os	2,6,8-Trimethyl-3-ethylchromone	_	,	112
		Ethylα-ethylacetoacetate Ethylα-benzylacetoacetate		POCI; H ₂ SO ₄	2,6,8-Trimethyl-3-ethylchromone	49		161
		Entily i descrity lacewacedak		(86%)	4.6.8-Trimethyl-3-benzylcoumarin			
		Ethyl benzoylacetate		H ₂ SO ₄	4-Phenyl-6,8-dimethylcoumarin	49	J	161
				(86%)	- 2 monte of dimension dimension			
3,5-Xyle	nol	Ethyl acetoacetate		H2SO4	4,5,7-Trimethylcoumarin	32-		5, 95
(3,5-đ		Ethyl a-methylacetoaceta	te	H2SO4	3,4,5,7-Tetramethylcoumarin	9-1	1 2	5,162
methy		Ethyl a-methylacetoaceta	te	P205	2,3,5,7-Tetramethylchromone	-		163
pheno		17.0		TT 00				25
2,5-Xyl (2,5-		Malic acid Ethyl acetoacetate		H ₂ SO ₄	5.8-Dimethylcoumarin	-	•	112
meth		Ethyl a-methylacetoscet	140	P ₂ O ₅ ;	2,5,8-Trimethylchromone		_ 1	12,160
phen		Linji u-metnyjaoeoospet	ate	POCI	2,3,5,8-Tetramethylchromone			
•		Ethyl a-ethylacetoacetat	e	P2O5;	2,5,8-Trimethyl-3-ethylchromone	_	_	112
				POC				
		Ethyl α-benzylacetoscet	ate	P2O6;	2,5,8-Trimethyl-3-benzylchromone	-	_	112
				POC	l ₂			
		Ethyl benzoylacetate		P2O5;	6.8-Dimethylflavone			112
Thyn	no!	Malie acid		P00	_	_		70
	scroj no	Ethyl acetoacetate		H ₂ SO			oor	39 164
	uw of	Ethyl a-methylacetoac		P ₂ O ₅ P ₂ O ₅	2.8-Dimethyl-5-isopropylchromone		_	164
4-Cb	loro-	Ethyl acetoacetate	· Lave	H ₂ SO	2.3.8-Trimethyl-5-isopropylchromo	ne	35	95
3,	5-di-	•		11200	6-Chloro-2,5,7-trimethylchromone		99	•••
	ethyl-							
	loasd							
	5-Trj-	Ethyl acetoscetate		P2O:	2,5,7,8-Tetramethylchromone		_	165
	etbyl- bezol							
	Jumeno.	Malic add		TT -				
, ,		Ethyl acetoscetate		H ₂ S H ₂ S			40	25 25
		Ethyl a-methylaceto:	cetat	H ₂ S			12	25 25
	Note: R	eferences 142-244 are listed			04 3,4,5,8,8-Pentamethylcoumarin	•	Poor	20
			27.0	p. 01–08.				

TABLE II
Condensations with Dihydric Phenols

		Condensing		Yield	Refer-
Phenol	Acid or Ester	Agent	Product	%	ence
Catechol	Acetonedicarboxylic ncid	H_2SO_4	8-Hydroxycoumarin-4-acetic acid	Poor	26
Guaiacol	Ethyl α-methylacetoacetate	P_2O_5	8-Methoxy-2,3-dimethy lchromone	5	166
Resorcinol	Diethyl malonate	C ₂ H ₅ ONa	Ethyl 7-hydroxycoumarin-4-ace- tate *	_	26
	Malio acid	H_2SO_4	7-Hydroxycoumarin	43-50	1, 8, 132
	Ethyl α -phonylformylacetate	P_2O_5	7-Hydroxy-3-phenylcoumarin		167
	Ethyl α -phenylformylacetate	ZnCl ₂	7-Hydroxy-3-phenylcoumarin	Poor	105
	Ethyl acetoacetate	H_2SO_4	7-Hydroxy-4-methylcoumarin	82-90	2, 133
	Ethyl acetoacetate	H ₂ SO ₄ (73%)	7-Hydroxy-4-methylcoumarin	-	168
	Ethyl acetoacetate	H ₂ SO ₄ (75%)	7-Hydroxy-4-methylcoumarin	96	169
	Ethyl acetoacetate	P_2O_{δ}	7-Hydroxy-4-methylcoumarin	63	101
	Ethyl acetoacetate	H ₃ PO ₄	7-Hydroxy-4-methylcoumarin	80	127
	Ethyl acetoacetate	$HCl + ZnCl_2$	7-Hydroxy-4-methylcoumarin	94	125
	Ethyl acetoacetate	HCl	7-Hydroxy-4-methylcoumarin	97	123
	Ethyl acetoacetate	FeCl ₃	7-Hydroxy-4-methylcoumarin	57	128
	Ethyl acetoacetate	SnCl ₄	7-Hydroxy-4-methylcoumarin	Quant.	128
	Ethyl acetoacetate	TiCl ₄	7-Hydroxy-4-methylcoumarin	- .	128
	Ethyl acetoacetate	C ₂ H ₅ ONa	7-Hydroxy-4-methylcoumarin	54	127
	Ethyl acetoacetate	CH ₃ CO ₂ Na	7-Hydroxy-4-methylcoumarin	72	127
	Ethyl acetoacetate	Boric anhy- dride	7-Hydroxy-4-methylcoumarin	50	127
	Ethyl acetoacetate (2 or more moles)	H ₂ SO ₄	Dimethyldicoumarin	10	170
	Ethyl acetoacetate (2 moles)	HCI	4,4'-Dimethylcoumarino-7,8,α-py- rone	20	62
	Ethyl α-chloroacetoacetate	H ₂ SO ₄	7-Hydroxy-3-chloro-4-methylcou- marin	-	32
	Ethyl α -chloroacetoacetate	P_2O_5	7-Hydroxy-3-chloro-4-methylcou- marin		109
	Methyl α-methylacetoacetate	H ₂ SO ₄	7-Hydroxy-3,4-dimethylcoumarin	_	2
	Ethyl a-methylacetoacetate	P ₂ O ₅	7-Hydroxy-3,4-dimethylcoumarin †	_	101, 109
	Ethyl α -methylacetoacetate	H ₃ PO ₄ ; CH ₃ CO ₂ Na; C ₂ H ₅ ONa	7-Hydroxy-3,4-dimethylcoumarin	_	109, 127
	Ethyl α -ethylacetoacetate	H ₂ SO ₄	7-Hydroxy-3-ethyl-4-methylcou- marin	_	109
	Ethyl α-ethylacetoacetate	H ₂ SO ₄ (73%)	7-Hydroxy-3-ethyl-4-methylcou- marin	54	101
	Ethyl α -ethylacetoacetate	P_2O_5	7-Hydroxy-3-ethyl-4-methylcou- marin	43	101, 109
	Ethyl α -propylacetoacetate	H_2SO_4 ; P_2O_5	7-Hydroxy-3-propyl-4-methylcou- marin	_	109
	Ethyl α -isopropylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-3-isopropyl-4-methyl- coumarin		109
	Ethyl α -hutylacetoacetate	H ₂ SO ₄ (80%)	7-Hydroxy-3-hutyl-4-methylcou- marin	_	47
	Ethyl α -isohutylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-3-isobutyl-4-methyl- coumarin	_	109
	Ethyl α -allylacetoacetate	H ₂ SO ₄	7-Hydroxy-3-allyl-4-methylcou- marin	97	70
	Ethyl α -allylacetoacetate	HCI	7-Hydroxy-3-chloropropyl- 4-methylcoumaria	87	70

Note: References 142-244 are listed on pp. 57-58.

^{*} The formation of this product was explained by the intermediate formation of acetonetricarhoxylic acid.

[†] Simonis and Remmert (ref. 5) carried out this condensation and assigned a chromone structure to the condensation product. Canter, Curd, and Robertson (ref. 101) have shown that the product is a coumarin derivative.

Phenol Resorciool (Cont'd)

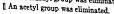
TABLE II-Continued

CONDENSATIONS WITH DIHYDRIC PHENOLS

		Condensing			Yield I	Refer-
	Acid or Ester	Agent		Product	%	ence
	Ethyl α-(α-hydroxy-β,β,β-tri-	H ₂ SO ₄ (78%)		lroxy-3-(a-hydroxy-8,8,8-tri-	12	72
	chloroethyl)acetoacetate Ethyl α-(α-hydroxy-β,β,β-tri-	P20s		oroethyl)-4-methylcoumarin droxy-3-(a-hydroxy-6,6,6-ki-	Poor	72
	ehloroethyl)acetoacetate Ethyl α-(α-hydroxy-β.β.β-trī-	POCl ₂		oroethyl)-4-methylcoumarin droxy-3-(a-hydroxy-8,8,8-tri-	36	72
	chloroethyl)acetoacetate Ethyl α-phenylacetoacetate	H ₂ SO ₄	chl	oroethyl)-4-methylcoumarin droxy-3-phenyl-4-methylcou-		105
	Etnyl a-phenylacetoacetate	H2304		uin		109
	Ethyl α-phenylacetoacetate	P ₂ O ₅		rdroxy-3-phenyl-4-methylcou- arin	-	-
	Ethyl α-p-methoxyphenyl- acetoacetate	H2SO4		ydroxy-3-p-methoxyphenyl-4- ethylcoumarin	-	17 i
	Ethyl a-benzylacetoacetate	H ₂ SO ₄	7-H	ydroxy-3-benzyl-4-methylcou-	55-65	105
	Ethyl α-benzylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅ ; H ₃ PO ₄ ; CH ₃ CO ₂ N ₂ ;	; 7-H	arin ydroxy-3-benzyl-4-methylcou- arin	-	109, 127
		C ₂ H ₅ ONa				470
	Ethyl a-benzylacetoacetate	POC12		lydroxy-3-bensyl-4-methylcou- narin	_	172
	Ethyl α-o-carboxybenzyl- acetoacetate	HCI		Iydroxy-3-o-carboxybensyl- i-methyl coumarin	_	79
	Ethyl acetocyanoacetate	H ₂ SO ₄		Hydroxy-4-methylcoumarin ‡	_	78
	Diethyl acetylmalonate	H2SO4		Hydroxy-4-methylcoumarin §	_	32, 104
	Diethyl acetosuceinate	H2SO4	Έt	hyl 7-hydroxy-4-methylcou- marin-3-acetate	30-63	75, 76
	Diethyl acetosuccinate	P2O5		thyl 7-hydroxy-4-methylcou- marin-3-acetate	Low	34, 127
	Diethyl acetosuccinate	H ₃ PO ₄	E	thyl 7-hydroxy-4-methylcou- marin-3-acctate	_	127
	Diethyl acetosuceinate	POCl ₃	E	thyl 7-hydroxy-4-methylcou-	Quant	. 34
	Diethyl acetosuccinate	AlCl ₃	7	marin-3-acetate -Hydroxy-4-methylcoumarin-	Quant	. 34
	Diethyl α-acetoglutarate	H ₂ SO ₄	F	3-acetic acid thyl 7-hydroxy-4-methylcou-	65	77
			7	marin-3-propionate '-Hydroxy-4-methylcoumarin-	6	
	Diethyl α-acetoglutarate	P2O5	7	3-propionic acid 7-Hydroxy-4-methyleoumarin-	_	173
	Diethyl α-acetoglutarate	H2PO4		3-propionic acid Ethyl 7-hydroxy-4-methylcou-	_	173
				marin-3-propionate 7-Hydroxy-4-methyleoumarin-		
	Diethyl a-acetoglutarate	VICI2		3-propionic acid 7-Hydroxy-4-methylcoumarin-	74	173
	Ethyl diacetylacetate	H2SO4		3-propionie acid		32
	Ethyl benzoylacetoaceta	te H-SO7	ZnCla	7-Hydroxy-4-methylcoumarin	_	32, 104
	Ethyl benzoylacetoaceta	te HC1	-4017	7-Hydroxy-4-phenylcoumarin	_	32, 104 174
	Ethyl phthalylacetoacet	ate HCl		7-Hydroxy-4-phenylcoumarin 7-Hydroxy-4-methylcoumarin-	_	79
	Diethyl acetonedicarbox	ylate H2SO4		3-benzoyl-o-carboxylie acid 7-Hydroxycoumarin-4-acetic aci	d 40	82,151
9	. ILPIPIPIPANO 140_044 11					

Note: References 142-244 are listed on pp. 57-58. ‡ The cyano group was eliminated.

A carbethoxyl group was eliminated.



CONDENSATIONS WITH DIHYDRIC PHENOLS

Acid or Ester	Condensing Ageot	Product	Yie %	
Acetonedicarboxylic acid	P ₂ O ₈	7-Hydroxycoumarin-4-acetic acid Dilactone of β,β-di(2,4-diliydroxy	2:	120
Acetonedicarboxylic ocid	POCl ₂	phenyl)glutario acid 7-Hydroxycoumarin-4-acetic acid	37	
		Dilactone of β,β-di(2,4-dillydroxy phenyl)glutnrio neid		
Acetooedicarboxylic acid	AlCl ₃	7-Hydroxycoumarin-4-acetic ocid Dilactoce of β,β-di(2,4-dilaydroxy pheoyl)glutaric acid		
Acetooedicarboxylic acid	SOC12	7-Hydroxycoumarin-4-acetic acid Dilactoce of β,β-di(2,4-dihydroxy	14 - 22	
Ethyl a-p-methoxyphonyl-	H ₂ SO ₄	phenyl)glutaric acid 7-Hydroxy-3-p-methoxyphenyl-		171
propionoacetate Ethyl butyroacetate	H ₂ SO ₄ (75%)	4-ethylcoumarin 7-Hydroxy-4-propylcoumarin	_	35
Ethyl α-p-methoxyphenyl- hutyroacetate	H ₂ SO ₄	7-Hydroxy-3-p-methoxyphenyl- 4-propyleoumarin	_	171
Ethyl α-p-methoxyphenyl- isovaleroacetate	H ₂ SO ₄	7-Hydroxy-3-p-methoxyphenyl- 4-isobutylcoumarin	-	171
Ethyl a-p-methoxypheoyl- caproylacetate	H ₂ SO ₄	7-Hydroxy-3-p-methoxyphenyl- 4-amylcoumarin	~	171
Ethyl benzoylacetate	H_2SO_4	7-Hydroxy-4-phenylcoumarin		2, 12
Ethyl benzoylacetate	H ₃ PO ₄	7-Hydroxy-4-phenylcoumario	•••	127
Ethyl benzoylacetate	HCl	7-Hydroxy-4-pheoylcoumarin	02	123
Ethyl a-benzylbeozoylacetate		7-Hydroxy-3-benzyl-4-phenyl- coumario	80	105
Ethyl α-benzylbenzoylacetate		7-Hydroxy-3-benzyl-4-phenyl- coumario	Poor	105
Diethyl benzoylsuccinate	H ₂ SO ₄ (85%)	marin-3-acetate	43	A7
Ethyl \gamma-phenylacetoacetate	H ₂ SO ₄	7-Hydroxy-4-beozylcoumarin ¶		80, frs
Ethyl δ-phenyl-β-ketoval- erate	H ₂ SO ₄	7-Hydroxy-4-(pheoethyl)cou- marin		17%
Ethyl veratroylacetate	H ₂ SO ₄	7-Hydroxy-4-veratrylcoumarin	-	hh,
Ethyl veratroylacetate	HCl	7-Hydroxy-4-veratrylcoumarin	90	65
Ethyl trimethylgalloylacetate	H ₂ SO ₄ (73%)	7-Hydroxy-4-(3,4,5-trimethoxy-phenyl)coumarin	A.O.,	174
Diethyl veratroylsuccinate	H ₂ SO ₄ (84%)	Ethyl 7-hydroxy-4-veratrylcou- marin-3-acetate	***	£7
Diethyl oxalacetate	C ₂ H ₅ ONa	Ethyl 7-hydroxycoumarin-4-car- boxylate	38-48	kk
Dimethyl oxalacetate	CH ₃ ONa	Metbyl 7-hydroxycoumarin-4-car- boxylate	-	特
Ethyl cyclopeotanone-2-car- boxylate	H ₂ SO ₄	7-Hydroxycyclopenteno-(1',2',4,3)- coumarin	1/1	91
Ethyl 4-methylcyclopenta- none-2-carboxylate	H ₂ SO ₄	7-Hydroxy-4'-metbylcyclopenteno- (1',2',4,3)-coumarin		11, 92
Ethyl cyclohexanone-2-car- boxylate	H ₂ SO ₄	Committee	Giran, g	14, 95
Etbyl cyclobexanone-2-car- boxylate	POCl ₃	7-Hydroxycyclohexeno-(1',2',4',3)- coumarin	3-10	124

Note: References 142-244 are listed on pp. 57-58.

Phenol

Resorciool

(Con!'d)

Note: References 142-244 are listed on pp. 20-20.

Baker and Robinson (ref. 106) reported the preparation of this compound by the Pechmann condensation of this compound by the Pechmann condensation of the Pechmann conde T Baker and Robinson (ref. 106) reported the preparation of the Sound Stevenson, and Thorge, i. Chem. Rep., 122, 211 (2015). The ather can be preparation of the sound Stevenson and Thorge, i. Chem. Rep., 122, 211 (2015). The ather can be preparation of the sound stevenson and Thorge, i. Chem. Rep., 122, 211 (2015). cinol with the material described as etdyl γ -purely activated and Litten (ref. 80) to be etdyl α -phenylardigues, then, then, 123, 1762 (1923). This material was later found by Sonn and Litten (ref. 80) to be etdyl α -phenylardigues, Therefore, Ther their condensation product with resorcinol is 7-hydroxy-3-phenyl-4-methylcoumarin.

CONDENSATIONS WITH DHIYDRIC PHENOLS

			01			Yield.	Refer-
5 11 1		1 11 Pater	Conden		Proluct	27	6300
Phenol Resorcinol	Ethyl	Acid or Ester 4-methylcyclohexa-	a 13 A 408:11		iroxy-4'-methyleyelobexeno-	_	97
(Cont'd)	100	ie-2-carboxylate	11:50,		,2(,4,3)-coumarin droxy-3'-methy leyelobesenoe	_	9, 97
	סמ	ne-2-carboxylate		(1'	(2',43)-mumarin		97
	no	ne-2-carboxylate	POCI:	(1	droxy-5'-methyleyelokexeno- (2',4,3)-coumarin		97
	-	d 6-methylcyclohexa- me-2-carboxylate	POC1	(1	rdroxy-6'-methylcyclohexeno- 1,2',4,3)-coumarin		E4
		Hydrindone-2-earboxylie iid	HC1	7-H	rdrexp-4,3-ied-nocoumaria	10	
		yl trans-β-decalone- -carboxylate	11:804		ydroxy-trans-octalino-(2',3',4,3)- oumarin	-	\$ 7
	Eth	yl indane-1,3-dione-2-car-	HCI	7-19	lydroxy-1'-ketoind-no-(2',3',3,4)-	7	83
	Et	oxylate hylβ-coumaranone-2-car-	H2SO	(85%) 7-1	oumarin lydroxyroumarono-(2',3',3,4)-	26	100
	Et	ooxylate hyl 5-methyl-β-coumara-	HCI	7-1	ounarin Iydroxy-5'-methykoumsrono-	ప	100
		none-2-carboxylate hyl 7-methyl-8-coumara-	11:50	4 (85%) 7-1	(2°,3°,3,4)-coumarin Hydroxy-7°-methylcoumarono-	23	100
	E	none-2-carboxylate thyl 6-methoxy-β-coumara	- 11C1		(2',3',3,4)-coumarin Hydroxy-6'-methoxy-coumarono-		100
	E	none-2-carboxylate thyl chroman-3-one-4-car-	1101	7-	(2',3',3,4)-coumarin Hydroxychromeno-(3',4',4,3)-	13	3 09
	I	hoxylate thyl 3-hydroxy-7-methoxy		7-	coumarin ·Hydroxy-7'-methoxychromeno-	_	. 99
	1	3-chromene-4-carboxylate Ethyl 3-hydroxy-8-methoxy	- HCI		(3',4',4,3)-coumarin -Hydroxy-8'-methoxychromeno-	_	. 63
		3-chromene-4-carboxylate Ethyl 3-hydroxy-6,7-dimetl oxy-3-chromene-4-car-		0, (82%) 7 5%)	(3',4',4,3)-coumarin '-Hydroxy-6',7'-dimethoxychro- meno-(3',4',4,3)-coumarin	1	1 2
		boxylate Ethyl 3-hydroxy-6,7-dimet oxy-3-chromene-4-car- boxylate	h- HC	١ :	7-Hydroxy-6',7'-dimethoxychro- meno-(3',4',4,3)-coumarin		9 59
		Methyl 3-hydroxyindolc- 2-carboxylate	H_2	SO ₄ (90%)	7-Hydroxyindolo-(2',3',3,4)-cou-		18 100
Reso	reinol	Malic acid	H.	SO ₄	marin 7-Methoxycoumarin	0	ant. 132
m	ono-	Ethyl acetoacetato		SO4; P ₂ O ₅		Qu	130
	ethyl	Acetonedicarhoxylic acid		SO4	7-Methoxy-4-methylcoumarin		26
et	her	Ethyl henzoylacetate		SO.	7-Methoxycoumarin-4-acetic acid		84
		Ethyl veratroylacetate		SO.	7-Methoxy-4-phenylcoumarin		86
_				,	7-Methoxy-4-(3',4'-dimethoxy-		_ 00
D	orcinol nonohutyl	Ethyl cyclohexanone-2-c boxylate	ar- P	OC13	phenyl)coumarin 3-Butoxy-7,8,9,10-tetrahydro-		157
	ther				6-dibenzopyrono		
	sorcinol	Ethyl acctoacetate	1	I2SO4	736.0		130
	dimethyl ether	Ethyl acetoacetate	I	H ₂ SO ₄ (80%; 87%)	7-Methoxy-4-methylcoumarin * '7-Methoxy-4-methylcoumarin *	•	13
	<i>~</i> .	Ethyl α-methylacetoace	tate 1	H ₂ SO ₄ (85%)	73(-1)		13
	Chloro-	Malic acid		H ₂ SO ₄		יי מו	_ ::
	resorcinol	Ethyl acetoacetate		H ₂ SO ₄	7-Hydroxy-6-chlorocoumarin 7-Hydroxy-4-methyl-6-chloroco	u-	25 41 26 41
		Ethyl acetoacetate		P ₂ O ₅	marin 7-Hydroxy-4-methyl-6-chloroco marin	u-	4
	Note: Ref	erences 142-244 are listed o		F.0	211111111		

^{••} Partial demethylation took place before the condensation.

CONDENSATIONS WITH DIHYDRIC PHENOLS

		Condensing		Yie.	ld Refer-
Phenol	Acid or Ester	Agent	Product	%	
4-Chloro- resorcinol	Ethyl α-chloroacetoacetate	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-3,6-dichloro-4-methyl- coumarin	_	41
(Cont'd)	Ethyl α-methylacctoacetato	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-3,4-dimethyl-6-chloro- coumarin	_	41
	Ethyl α -ethylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅		-	41
	Ethyl α-propylacctoacetate	H ₂ SO ₄ ; P ₂ O ₅		_	41
	Ethyl $lpha$ -isobutylacetoacetstc	H ₂ SO ₄	7-Hydroxy-3-isohutyl-4-methyl- 6-chlorocoumarin	_	41
	Ethyl α-benzylacetoacetate	H ₂ SO ₄	7-Hydroxy-3-benzyl-4-methyl- 6-chlorocoumarin	_	41
	Diethyl acetosuccinate	H_2SO_4	Ethyl 7-hydroxy-4-methyl-6-chloro- coumarin-3-acetate	-	41, 42
	Diethyl acetosuccinate	POCl ₃	Ethyl 7-hydroxy-4-methyl-6-chloro- coumarin-3-acetate	-	42
	Acctonedicarboxylio acid	H ₂ SO ₄	7-Hydroxy-6-chlorocoumarin- 4-acetic acid	_	41
	Ethyl benzoylacetate	H ₂ SO ₄	7-Hydroxy-4-phenyl-6-chlorocou- marin	-	41
4-Bromo- resorcinol	Ethyl acetoacetate	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-4-methyl-6-hromocou- marin	_	43, 177
	Ethyl α-methylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-3,4-dimethyl-6-hromo- coumarin	-	43
	Ethyl α-ethylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅	7-Hydroxy-3-ethyl-4-methyl- 6-hromocoumarin	-	43
	Diethyl acetosuccinate	POC13	Ethyl 7-hydroxy-4-methylcou- marin-3-acctate	_	42
2-Nitro- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-nitrocou- marin	60	41
	Ethyl α-methylacetoacetate	H ₂ SO ₄	7-Hydroxy-3,4-dimethyl-8-nitro- coumarin	15	41
4-Nitro- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-6-nitrocou- marin	_	44
	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-6-nitrocou- marin	3	118
2-Amino- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-aminocou- marin	-	177
2-Methyl-	Malic acid	H ₂ SO ₄	7-Hydroxy-8-methylcoumarin	_	178
resorcinol	Ethyl acetoacetate Ethyl benzoylacetate	H ₂ SO ₄ H ₂ SO ₄	7-Hydroxy-4,8-dimethylcoumarin 7-Hydroxy-4-phenyl-8-methylcou- marin	89	62 179
4-Methyl-	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4,6-dimethylcoumarin	Quant.	180
resorcinol	Diethyl acetosuccinate	POCl ₃	Ethyl 7-hydroxy-4,6-dimethylcou- marin-3-acetate	_	181
5-Methyl-	Malic acid	H_2SO_4	7-Hydroxy-5-methylcoumarin	Good	39, 40
resorcinol	Ethyl acetoacetate	H ₂ SO ₄	5-Hydroxy-4,7-dimethylcoumarin ††	91	31
(orcinol)	Ethyl acetoacetate	H ₂ SO ₄ (73%)	5-Hydroxy-4,7-dimethylcoumarin ††	68	168
	Ethyl acetoacetate	P ₂ O ₅	5-Hydroxy-4.7-dimethylcoumarin	-	33
	Ethyl acetoacetate	H ₃ PO ₄ (coned. and 85%)	5-Hydroxy-4,7-dimethylcoumarin	55	127, 182

^{††} Müller (ref. 151) who also carried out these condensations, assigned the 7-hydroxycoumarin structure to the product. This is incorrect as the product was shown earlier, by Collie and Chrystall, J. Chem. Soc., 91, 1804 (1907), to have the 5-hydroxycoumarin structure.

Phenol

5-Methylresorcinol
(orcinol)
(Cont'd)

TABLE II-Continued

CONDENSATIONS WITH DIHYDRIC PHENOLS

		Condensing		Yield	Refer-
	Acid or Ester	Agent	Product	%	ence
	Ethyl α -chloroacetoacetate	H2SO4	5-Hydroxy-3-chloro-4,7-dimethyl- coumarin	60	32
	Ethyl α-chloroacetoacetate	P204	5-Hydroxy-3-chloro-4,7-dimethyl- coumarin		33
	Ethyl a-methylacetoacetate	P2O5	5-Hydroxy-3,4,7-trimethylcoumarin		33
	Ethyl α-ethylacetoacetate	H ₂ SO ₄	5-Hydroxy-3-ethyl-4,7-dimethyl- coumarin		33
	Ethyl α-butylacetoacetate	H ₂ SO ₄	5-Hydroxy-3-butyl-4,7-dimethyl- coumarin		37
	Ethyl α-hutylacetoacetate	POCl ₃	5-Hydroxy-3-hutyl-4,7-dimethyl- coumarin	62	182
	Ethyl &-allylacetoacetate	HCl	5-Hydroxy-3 (β-chloropropyl)- 4,7-dimethylcoumarin	-	124
	Ethyl α-(α-hydroxy-β,β,β-tri- chloroethyl)acetoacetate	- POCl ₁	5-Hydroxy-3(α-hydroxy-β,β,β-tri- chloroethyl)-4,7-dimethylcou- marin	30	72
	Ethyl α-benzylacetoacetate	H ₂ SO ₄	7-Hydroxy-3-benzyl-4,5-dimethyl- coumnrin 11		105
	Diethyl acetosuccinate	H ₂ SO ₄ ; P ₂ O ₅ H ₂ PO ₄		_	34, 127
	Diethyl acetosuccinate	POCl ₂	Ethyl 5-hydroxy-4,7-dimethylcou- marin-3-acctate	67	34
	Diethyl α-acetoglutarate	H ₂ SO ₄	Ethyl 5-hydroxy-4,7-dimethylcou- marin-3-propionate and 5-hy- droxy-4,7-dimethylcoumarin- 3-propionic acid	-	77
	Diethyl a-scetoglutarate	PrOs	5-Hydroxy-4,7-dimethylcoumarin- 3-propionic acid	_	173
	Diethyl α-acetoglutarate	HCI	Ethyl 5-hydroxy-4,7-dimethylcou- marin-3-propionate and 5-hy- droxy-4,7-dimethylcoumarin- 3-propionio neid	-	77, 173
	Acetonedicarboxylic acid	H ₂ SO ₄	5-Hydroxy-7-methylcoumarin- 4-acetic acid	Good	26
	Citric acid	H_2SO_4	5-Hydroxy-7-methylcoumarin- 4-acetic acid and orcin-aurin	_	151
	Ethyl butyroacetate	H ₂ SO ₄ (75	%) 5-Hydroxy-4-propyl-7-methylcou- marin	. –	35
	Ethyl γ-phenylacetoacets				81
	Ethyl α-benzylhenzoylac		5-Hydroxy-3-benzyl-4-phenyl- 7-methylcoumarin §§	_	103
	Ethyl cyclopentanone-2- boxylate		5-Hydroxy-7-methyl-3,4-cyclo- pentenocoumarin	_	. 36
	Ethyl cyclopentanone-2- boxylate		5-Hydroxy-7-methylcyclopentene (1',2',4,3)-coumarin	o- 57	7 91
	Ethyl 4-methylcyclopen none-2-carboxylate	-	5-Hydroxy-7,4'-dimethyloyclo- penteno-(1',2',4,3)-coumarin	_	. 91
	Ethyl cyclohexanone-2- boxylate	car- H ₂ SO ₄	1-Hydroxy-3-methyl-7,8,9,10-tet hydro-8-dibenzopyrone	.га- 3	5 10
Vο	te: References 142-244 are listed of				

tt By analogy with the other compounds obtained in the condensation of orcinol with β -ketonic esters, this compound is probably a 5-bydroxycoumarin derivative.

^{§§} By analogy with the other compounds obtained in the condensation of orcinol with β -ketonic esters, this compound is probably a 5-hydroxy-coumarin derivative. The structure originally assigned (7-hydroxy-3-benryl-4-phenyl-5-methyl-coumarin) is incorrect; refs. 105, 106.

Condensations with Dihydric Phenols

Phenol	Acid or Ester	Condensing Agent	g Product	Yield	d Refer- ence
5-Methyl- resorcinol	Ethyl cyclohexanone-2-car- boxylate	POCl ₃	5-Hydroxy-7-methylcyclohexeno- (1',2',4,3)-coumarin	_	124
(orcinol) (Cont'd)	Ethyl cyclohexanone-2-car- hoxylate	POCl ₃	1-Hydroxy-3-methyl-7,8,9,10-tetra- hydro-6-dihenzopyrone	66	10
	Ethyl 4-methylcyclohex- anone-2-carboxylate	POCI ₃	5-Hydroxy-7,4'-dimethylcyclohex- eno-(1',2',4,3)-coumarin	-	97
	Ethyl 5-methylcyclohex- anone-2-carboxylate	H ₂ SO ₄ ; POCl ₃	5-Hydroxy-7,5'-dimethylcyclohex- eno-(1',2',4,3)-coumarin	-	97
	Ethyl 5-methylcyclohex- anone-2-carhoxylate	POCl ₃	1-Hydroxy-3,9-dimethyl-7,8,9,10- tetrahydro-6-dihenzopyrone	62	10
	Ethyl 6-methylcyclohex- anone-2-carboxylate	POCl ₃	5-Hydroxy-7,6'-dimethylcyclohex- eno-(1',2',4,3)-coumarin	-	97
	Ethyl trans-β-decalone-3-car- hoxylate	-	5-Hydroxy-7-methyl-trans-octalino- (2',3',4,3)-coumarin	_	97
2-Ethyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-ethylcou- marin	79	183
4-Ethyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-6-ethylcou- marin	49- Quant.	184, 185
	Ethyl acetoacetate	H ₂ SO ₄ (73%)	marin	80-85	186
	Ethyl α-methylacetoacetate	H ₂ SO ₄ (73%)	coumarin	90	55
	Ethyl α-methylacetoacetate	POCI ₃	7-Hydroxy-3,4-dimethyl-6-ethyl- coumarin	_	187
	Ethyl α-ethylacetoacetate	H ₂ SO ₄ (73%)	coumarin	75	55
	Ethyl α-ethylacetoacetate	POCl ₃	7-Hydroxy-3,6-diethyl-4-methyl- coumarin	_	187
	Ethyl α-propylacetoacetate	H ₂ SO ₄ (73%)	7-Hydroxy-3-propyl-4-methyl- 6-ethylcoumarin	65	55
	Ethyl α-propylacetoacetate	POCl ₃	7-Hydroxy-3-propyl-4-methyl- 6-ethylcoumarin	_	187
	Ethyl α-hutylacetoacetate	H ₂ SO ₄ (73%)	7-Hydroxy-3-hutyl-4-methyl- 6-ethylcoumarin	_	55
	Ethyl α-hutylacetoacetate	POCI ₃	7-Hydroxy-3-butyl-4-methyl- 6-ethylcoumarin	45	187
	Ethyl α-allylacetoacetate	H ₂ SO ₄ (73%)	7-Hydroxy-3-allyl-4-methyl- 6-ethylcoumarin		55
	Ethyl α -(α -hydroxy- β , β , β -tri- chloroethyl)acetoacetate	H ₂ SO ₄ (80%)	7-Hydroxy-3-(α-hydroxy-β,β,β-tri- chloroethyl)-4-methyl-6-ethyl coumarin	Poor	74
	Ethyl α-(α-hydroxy-β,β,β-tri- chloroethyl)acetoacetate	POCl ₃	7-Hydroxy-3-(α-hydroxy-β,β,β-tri- chloroethyl)-4-methyl-6-ethyl- coumarin	27	74
	Diethyl acetosuccinate	H ₂ SO ₄ (80%)	Ethyl 7-hydroxy-4-methyl-6-ethyl- coumarin-3-acetate		181
	Diethyl acetosuccinate	POCl ₃	Ethyl 7-hydroxy-4-methyl-6-ethyl- coumarin-3-acetate		181
	Ethyl benzoylacetate		7-Hydroxy-4-phenyl-6-ethylcou- marin	90	55
	Ethyl cyclopentanone-2-car- bexylate	H ₂ SO ₄	7-Hydroxy-6-ethylcyclopenteno- (1',2',4,3)-coumarin	32	91

III Sen and Basu (ref. 94) have carried out the same condensation and assigned the 7-hydroxy structure to the condensation product. Chowdhry and Desai (ref. 97) have shown this to be incorrect and have assigned the 5-hydroxy-coumarin structure.

CONDENSATIONS WITH DIHYDRIC PHENOLS

		Condensing		11010	Refer-
Phenol	Acid or Ester	Agent	Product	%	•
4-Ethyl- I resorcinol	Ethyl 4-methylcyclopenta- none-2-carhoxylate	H ₂ SO ₄	7-Hydroxy-4'-methyl-6-ethylcyclo- penteno-(1',2',4.3)-coumarin	_	91
(Cont'd)	Ethyl cyclohexanone-2-car- boxylate	POCl ₂	7-Hydraxy-6-ethylcyclohexeno- (1',2',4,3)-coumarin		124
,	Ethyl 4-methylcyclohex- anone-2-carboxylate	H2SO4	7-Hydroxy-4'-methyl-6-ethylcyclo- hexeno-(1',2',4,3)-coumarin		96
	Ethyl 5-methylcyclohex- anone-2-carboxylate	H ₂ SO ₄	7-Hydroxy-5'-methyl-6-ethylcyclo- hexeno-(1',2',4,3)-coumarin		96
	Ethyl 6-methylcyclohex- anone-2-carboxylate	POCI:	7-Hydroxy-6'-methyl-6-ethylcyclo- hexeno-(1',2',4,3)-cnumarin		97
	Ethyl trans-β-decalone-3-car- boxylate	H ₂ SO ₄	7-Hydrnxy-6-ethyl-trans-octalino- (2',3',4,3)-coumarin	_	96
5-Ethyl- resorcinol	Ethyl 1-methylcyclohexan- 3-one-4-carboxylate	H ₂ SO ₄	5-Hydroxy-5'-methyl-7-ethyl- 3,4-cyclohexenocnumarin	_	37
4-Propyl- resorcinol	Ethyl acetnacetate	H2SO4	7-Hydroxy-4-methyl-6-prapyl- coumarin	_	185
	Ethyl α-(α-hydroxy-β,β,β- trichloroethyl)acetoacetate	POCt ₃	7-Hydroxy-3-(α-hydroxy-β,β,β-tri- chloroethyl)-4-methyl-6-propyl- coumarln	Low	74
	Diethyl acetosuccinate	H ₂ SO ₄	Ethyl 7-hydroxy-4-methyl-6-propyl- coumsrin-3-acetate	38	181
	Diethyl acetosuccinate	POCl ₁	Ethyl 7-hydroxy-4-methyl-6-prapyl- coumarin-3-acetate	Quant.	
5-Propyl- resorcinol	Ethyl 5-methylcyclohex- anone-2-carboxylate	POCl ₃	1-Hydroxy-3-propyl-9-methyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	55	38
4-Butyl- resorcinol	Ethyl α-(α-hydroxy-β,β,β- trichloroethyl)acetoacetate	POCl ₃	7-Hydroxy-3-(\alpha-hydrnxy-\beta,\beta,\beta-tri- chloroethyl)-4-methyl-6-hutyl- coumarin	_	168
	Diethyl acetosuccinate	POCl ₃	Ethyl 7-hydroxy-4-methyl-6-hutyl- coumsin-3-acetate	-	181
5-Butyl- resorcinol	mione-2-cut boxytate	POC1 ₃	1-Hydroxy-3-hutyl-9-methyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	59	38
2-Isoamyl- resorcino	Malic acid	H_2SO_4	7-Hydroxy-8-isoamylcoumarin	39	189
(tetra- hydro-	zenya accoracetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-isoamyl- coumarin	20	169
tuhanol)		· n	7-Hydroxy-7'-methoxy-8-isoamyl- chromeno-(3',4',4,3)-coumarin	_	99
	Ethyl 3-hydroxy-8-methox 3-chromene-4-carboxylar	ło.	7-Hydroxy-8'-methoxy-8-isoamyl- chromeno-(3',4',4,3)-coumarin	49	99
9.71	Ethyl 3-hydroxy-6,7-dime oxy-3-chromene-4-car- boxylate	th- H ₂ SO ₄ (85)	%) 7-Hydroxy-6',7'-dimethoxy-8-iso- amylchromeno-(3',4',4,3)-cou- marin		99
2-Isoamyl resorcin mono- methyl ether 4-Isoamy	ol .	H ₂ SO ₄	7-Methoxy-8-isoamylcoumarin	66	190, 191
resorci		H ₂ SO ₄ ; H ₂ SO ₄ ; POCl ₃ ; AlCl ₃	7-Hydroxy-6-isoamylcoumarin 7-Hydroxy-4-methyl-6-isoamyl- coumarin	-	. 192 . 30

CONDENSATIONS WITH DIHYDRIC PHENOLS

		Condensing		Yiel	a be
Phenol	Acid or Ester	Agent	Product	50	d Refer- ence
5-Amyl- resorcinol	Ethyl acetoacetate	H:50.	5-Hydroxy-4-methyl-7-amylcou- marin	****	30
(olivetol)	Ethyl acetoacetate	POCI ₂	6-Hydroxy-f-methyl-7-amylcou- marin	85	192, 193
	Ethyl a-butylacetoacetate	POCI:	5-Hydroxy-3-butyl-4-methyl- 7-amylcoumarin	66	182, 193
	Ethyl cyclopentanone-2-car- boxylate	112504	5-Hydroxy-7-amyl-3,4-cyclopen- tenocoumarin		30
	Ethyl cyclohexanone-2-car- boxylate	112SO4	5-Hydroxy-7-amyl-3,4-cyclohexeno coumarin		36
	Ethyl cyclohexanone-2-car- boxylate	POCI ₂	1-Hydroxy-3-amyl-7,8,9,10-tetra- hydro-6-dibenzopyrone	82	93
	Ethyl 4-methylcyclolicxa- none-2-carboxylate	POCl ₃	I-Hydroxy-3-amyl-8-methyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	76	93
	Ethyl 1-methylcyclohexan- 3-one-4-carboxylate	II2SO4	5-Hydroxy-5'-methyl-7-amyl- 3,4-cyclohexenocoumnin	91	9
	Ethyl 5-methylcyclohexa- none-2-carboxylate	POCI3	I-Hydroxy-3-amyl-9-methyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	57-75	93, 194
	Ethyl 5-ethyleyelohexanone- 2-carboxylate	POCI ₃	1-Hydroxy-3-amyl-9-ethyl-7,8,9,10- tetrahydro-6-dibenzopyrone	46	98
	Ethyl 6-methylcyclohexa- none-2-carlioxylate	POCI ₃	I-Hydroxy-3-amyl-10-methyl- 7,8,9,10-tetrnhydro-6-dibenzo- pyrone		93
	Ethyl 3,5-dimethylcyclohexa- none-2-carboxylate	POCI3	1-Hydroxy-3-amyl-7,9-dimethyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	દર	98
	Ethyl 4,5-dimethylcyclohexa- none-2-carboxylato	POCI3	1-Hydroxy-3-amyl-8,9-dimethyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	61	98
	Ethyl 5,5-dimethylcyclohexa- none-2-carboxylate	POCI ₃	1-Hydroxy-3-amyl-9,9-dimethyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	33	98
	Ethyl cycloheptanone-2-car- boxylate	POCI ₃	5-Hydroxy-7-amyl-3,4-penta- methylenecoumarin	45	98
5-Isoamyl- resorcinol	Ethyl 1-methyleyclohexan- 3-one-4-carboxylate	H ₂ SO ₄	5-Hydroxy-5'-methyl-7-isoamyl- 3,4-eyelohexenocoumarin	-	37
4-Hexyl- resorcinol	Ethyl ncetoacetate	H ₂ SO ₄ (82%)	7-Hydroxy-4-methyl-6-hexyl- coumarin	39	195
5-Hexyl- resorcinol	Ethyl 5-methyleyelohexa- none-2-carboxylate	POCI ₃	1-Hydroxy-3-hexyl-9-methyl- 7,8,9,10-tetrahydro-6-dihenzo- pyrone	52	38
5-Isohexyl- resorcinol	Ethyl 1-methyleyclohexan- 3-one-4-carhoxylate	H ₂ SO ₄	5-Hydroxy-5'-methyl-7-isohexyl- 3,4-cyclohexenocoumarin	-	37
5-Heptyl- resorcinol	Ethyl 5-methyleyelohexa- none-2-carboxylate	POCI ₃	1-Hydroxy-3-heptyl-9-methyl- 7,8,9,10-tetrahydro-6-dihenzo- pyrone	59	38
5-Octyl- resoreinol	Ethyl 5-methylcyclohexa- none-2-carhoxylate	POCI ₃	1-Hydroxy-3-octyl-9-methyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone	59	38
4-Dodecyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄ ; POCl ₃ ; AlCl ₃	7-Hydroxy-4-methyl-6-dodecyl- coumarin		30

CONDENSATIONS WITH DIHYDRIC PHENOLS

Vield Refer-

		Condensing		Yield	Refer-
Phenol	Acid or Ester	Agent	Product	%	ence
l-Hexadecyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄ ; POCl ₁ ; AlCl ₂	7-Hydroxy-4-methyl-6-hexadecyl- coumarin	_	30
4-Octadecyl- resorcinol	Ethyl acetoacetate	POCI3	7-Hydroxy-4-methyl-6-octadecyl- coumarin	_	196
	Misc	ellaneous C-A	lkylresorcinols		
5-Alkyl- resorcinol	Ethyl 5-methylcyclohexa- none-2-carboxylate	POC13	1-Hydroxy-3-alkyl-9-methyl- 7,8,9,10-tetrahydro-6-dibenzo- pyrone CH ₃ OH		
			CO-O R = alkyl group		
5-Alkyl suhstituer	.4		3-Alkyl substituent		
1-Methyl-	Ethyl 5-methylcyclohexa-	Doo:		70	197
hutyl	none-2-carboxylate	POC13	1-Methylhutyl	70	10.
1-Ethylhut	yl Ethyl 5-methylcyclohexa- none-2-carboxylate	POC13	1-Ethylhutyl	73	197
1-Methyl- pentyl	Ethyl 5-methylcyclohexa- none-2-carboxylate	POCI3	1-Methylpentyl	53	197
1-n-Propyl pentyl	 Ethyl 5-methylcyclohexa- none-2-carboxylate 	POC13	1-n-Propylpentyl	51	197
1-Methyl- hexyl	Ethyl 5-methylcyclohexa-	POC13	1-Methylhexyl	47	197
1-Methyl-	Ethyl 5-methylcyclohera-	POCl ₃	1-Methylheptyl	62	197
heptyl —CH(CH	none-2-carboxylate		2-Metaly mepty i	-	
(CH ₂) ₅ —CH(CH	CH ₃ none-2-carboxylate	POCl ₃	-CH(CH ₃)(CH ₂) ₆ CH ₃	38	
(CH ₂);		POC13	-CH(CH ₃)(CH ₂) ₇ CH ₃	41	198
-CH ₂ Cl (CH ₃)	H- Ethyl 5-methylcyclohexa- CH ₂ - none-2-carboxylate	POCl ₃	-CH ₂ CH(CH ₃)CH ₂ CH ₂ CH ₃	60	198
CH ₂ C CH ₂ C CH(C CH ₂ C	H ₂ - Ethyl 5-methylcyclohexa- H ₃)- none-2-carhoxylate	POCI3	CH ₂ CH ₂ CH(CH ₃)CH ₂ CH ₃	7:	2 198
-CH ₂ C CH ₂ C (CH ₃	CH- none-2-carhoxylate	POCl ₃	-CH ₂ CH ₂ CH ₂ CH(CH ₃) ₂	7	3 198
C3H	none-2-carboxylate		-C(CH ₃) ₂ C ₃ H ₇	7	3 199
C(CI CH(i C₂H	CH ₃)- Ethyl 5-methylcyclohexa- CH ₃)- none-2-carboxylete	POCl ₃	$-C(CH_3)CH(CH_3)C_2H_5$	3	0 199
CH(C ₂ H ₅)- Ethyl 5-methylcyclohexa- CH ₃)- none-2-carboxylate	POCl ₃	CH(C ₂ H ₅)CH(CH ₃)CH ₃	2	8 199
D)D—	In none-2-carboxylate	•	-C(CH ₃) ₂ C ₆ H ₁₃	;	37 199
C# CH	(CH ₃)- Ethyl 5-methylcyclohexa (CH ₃)- none-2-carboxylate	- POCl ₃	—СН(СН ₃)СН(СН ₃)С ₅ Н ₁₁	:	24 199

CONDENSATIONS WITH DIHYDRIC PHENOLS

Phenol	4 * 2	Condensi	ng	Yield	Refer-
	Acid or Ester	Agent	Product	%	ence
β-Resorcylic acid	Molic acid	H ₂ SO ₄	7-Hydroxycoumarin-6-carboxylic acid	30	45,200, 201
	Malic ocid	H ₂ SO ₄	7-Hydroxycoumarin-6-carboxylic acid (isolated as methyl 7-meth	20	120
			oxycoumarin-6-carboxylate) 5-Hydroxycoumarin-6-carboxylic acid (isolated as methyl 5-meth- oxycoumarin-6-carboxylate)	1	
	Etbyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methylcoumarin- 6-carboxylic acid	21	45
	The same		7-Hydroxy-4-methylcoumarin	Traces	
	Ethyl ocetoacetate	AlCl ₃	5-Hydroxy-4-methylcoumario- 6-carboxylic acid	14	53
	Ethyl 4-methylcyclopenta- none-2-carboxylate	H ₂ SO ₄ (73%	6) 7-Hydroxy-6-carhoxy-3,4-(4'- methylcyclopenteno)coumarin	-	48
Mari	Ethyl cyclohexanone-2-car- hoxylate	H ₂ SO ₄ (739	6) 7-Hydroxy-6-carboxy-3,4-cyclo- hexenocoumarin	-	48
Methyl <i>β</i> - resorcylate	Molie acid	H_2SO_4	7-Hydroxycoumario-6-carboxylic acid	~	45
	Ethyl acetoacetate	H ₂ SO ₄ (80%		43	45
			7-Hydroxy-4-metbylcoumario- 6-carboxylic acid	31	
	Ethyl acetoacetate	P_2O_5	Methyl 7-bydroxy-4-methylcou- marin-6-carboxylate	3	45
	Ethyl acetoacetate	POCl ₃	Mcthyl 7-hydroxy-4-metbylcou- marin-6-carboxylate	5	45
	Ethyl acetoacetate	AlCl ₃	Methyl 5-bydroxy-4-methylcou- marin-6-carboxylate	18	53
			Methyl 7-bydroxy-4-methylcou- mario-6-carhoxylate	2	
	Etbyl acetoacetate	HCl	Methyl 7-bydroxy-4-methylcou- marin-6-carhoxylate	19	45
	Ethyl acetoacetate	$ZnCl_2$	Methyl 5-bydroxy-4-methylcou- marin-6-carboxylate		53
	•		Methyl 7-hydroxy-4-methylcou- marin-6-carhoxylate	_	
	Ethyl α-chloroacetoacetate	H ₂ SO ₄ (80%)	Methyl 7-hydroxy-3-chloro-4- methylcoumarin-6-carboxylate	6	46
	Etbyl α-metbylacetoacetate	H ₂ SO ₄ (80%)	Methyl 7-hydroxy-3,4-dimethyl- coumarin-6-carhoxylate	20	47
			7-Hydroxy-3,4-dimethylcoumarin- 6-carboxylic acid	7	
	Ethyl α -ethylacetoacetate	H ₂ SO ₄ (80%)	Methyl 7-hydroxy-3-ethyl-4- methylcoumarin-6-carboxylate	_	47
	Ethyl α -propylacetoacetate	H ₂ SO ₄ (80%)	Methyl 7-hydroxy-3-propyl-4- methylcoumarin-6-carboxylate 7-Hydroxy-3-propyl-4-methyl-	_	47
	The second second	TT SO. (8007)	coumarin-6-carboxylic acid Methyl 7-bydroxy-3-butyl-4-		17
	Ethyl α-hutylacetoacetate	H ₂ SO ₄ (80%)	methylcoumarin-6-carboxylate 7-Hydroxy-3-butyl-4-methyl-	- 4	17
	Ethyl α-henzylacetoacetate	H ₂ SO ₄ (80%)	coumarin-6-carboxylic acid Methyl 7-hydroxy-3-henzyl-4- methylcoumarin-6-carboxylate	- 4	7

Condensations with Dihydric Phenols

	CONDENSATIO	NS WITH D	IHIDRIC I HEROES		
		Condensing		1,014	Refer- ence
Phenol	Acid or Ester	Agent	Product	%	•
	Diethyl acetosuccinate	H ₂ SO ₄ (80%)	Methyl 7-hydroxy-4-methyl-	54	42
resorcylate	Dietaji acciosocianio	-	coumarin-6-carboxylate 🟋	_	48
	Ethyl α-benzoylacetoacetate	H ₂ SO ₄ (80%)	Methyl 7-hydroxy-4-pheoylcou-	6	40
(00.11 2)	,		marin-6-carboxylate	2	
			7-Hydroxy-4-phenylcoumarin-	2	
			6-carboxylic acid	8	46
	Diethyl acetonedicarboxylate	H ₂ SO ₄ (80%)	Ethyl 7-hydroxy-6-carbomethoxy-	·	
			coumarin-4-acctate	12	
			7-Hydroxy-6-carbomethoxycou-		
	TW -1 1 1 0	11 CO. (7207)	marin-4-acctio acid 7-Hydroxy-6-carbomethoxy-	42	48
	Ethyl eyclopentanone-2-car-	H ₂ SO ₄ (73%)	3,4-cyclopenteoocoumarin		
	boxylate Ethyl 4-methylcyclopenta-	H ₂ SO ₄ (73%)		46	48
	none-2-carboxylate	11201 (13/8)	(4'-methylcyclopenteno)coumarin		_
	Ethyl eyclohexanone-2-cat-	H ₂ SO ₄ (73%)		61	48
	boxylate	22200 (1070)	cyclohexenocoumarin		
	Ethyl cyclohexanone-2-car-	POCl ₂	7-Hydroxy-6-carbomethoxy-3,4-	77	48
	boxylate		cyclohexeoocoumario		48
	Ethyl cyclohexanone-2-car-	AlCl ₂	7-Hydroxy-6-carbomethoxy-3,4-	77	40
	boxylate		cyclohexenocoumarin		49
γ-Resorcylic	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methylcoumarin-	60	30
acid	Total and a second	** 50	8-carboxylic neid	46	17
2-Acetyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄ (78%		-10	
resorcinor	Ethyl acetoacetate	AlCla	coumario	74	17
	Minji acconcernie	AICI3	7-Hydroxy-4-methyl-8-acetylcou- marin	• • •	
	Ethyl acetoacetate	FeCl ₂	7-Hydroxy-4-methyl-8-acetylcou-		128
	•		marin		
	Diethyl acetosuccinate	H2SO4 (809			42
			marin-3-acetie ncid		42
	Diethyl acetosuccinate	FOCI3	Ethyl 7-hydroxy-4-methyl-8-acetyl	. –	42
4-Acetyl-	Malic acid		coumarin-3-acetate		202
resorcin		H ₂ SO ₄	7-Hydroxycoumarin *	40	
(resacet	Q - DOGGOGGGGGGGG	POC13	7-Hydroxy-4-methyl-6-acetyl-	41	J
phenon		POCl ₂	coumarin	40	12
	•	10013	7-Hydroxy-4-methyl-6-acetyl- coumarin	3,	•
			5-Hydroxy-4-methyl-6-acctyl-	_	-
	m., .		coumarin		
	Ethyl acetoacetate	AlCi2	5-Hydroxy-4-methyl-6-acetyl-	37-	41 53
	Ethyl - mathalast		coumarin		7 116
	Ethyl α-methylacetoace	etate AlCl3	5-Hydroxy-3.4-dimethyl-6-acetyl-		7 116
	Ethyl α-ethylacetoacet	ate AlCiz	eoumarin		116
			5-Hydroxy-3-ethyl-4-methyl- 6-acetylcoumarin	-	
	Ethyl α -benzylacetoac	etate AlCl3	5-Hydroxy-3-benzyl-4-methyl-		116
		=	6-acetylcoumarin		
	Ethyl cyclopentanone-	2-car- POCl3	7-Hydroxy-6-acetyl-3,4-cyclo-		25 48
	hoxylate Ethyl cyclopentanone	0 115	pentenocoumarin		
	boxylate	-2-car- AlCl ₃	5-Hydroxy-6-acetyl-3,4-cyclo-		48
	Ethyl 4-methylcyclop	enta- AlCla	pentenocoumarin		48
	none-2-carboxylate	- 111012	5-Hydroxy-6-acetyl-3,4-(4'-meth	yl-	10
Not	e: References 149-944 11 1		cyclopenteno)coumarin		

Note: References 142-244 are listed on pp. 57-58.
§¶ In this condensation a —CH₂CO₂C₂H₅ group was eliminated.
• In this condensation an acetyl group was eliminated.

CONDENSATIONS WITH DIHYDRIC PHENOLS

704		Condensing	3	Yielo	l Refer-
Phenol	Acid or Ester	Agent	Product	%	ence
4-Acetyl- resorcinol	Ethyl cyclohexanone-2-car- hoxylate	POCl ₃	7-Hydroxy-6-acetylcyclohexeno- (1',2',4,3)-coumarin	_	96
(resaceto- phenone)	Ethyl cyclohexanone-2-car- boxylate	AlCl ₃	5-Hydroxy-6-acetyl-3,4-cyclo- hexenocoumarin	82	48
(Cont'd)	Ethyl 4-methylcyclohexa- none-2-carhoxylate	POCl ₃	7-Hydroxy-4'-methyl-6-acetyl- cyclohexeno-(1',2',4,3)-coumarin	_	96
	Ethyl 5-methylcyclohexa- none-2-carhoxylate	POCl ₃	7-Hydroxy-5'-methyl-6-acetyl- cyclohexeno-(1',2',4,3)-coumarin	_	96
~	Ethyl trans-β-decalone- 3-carhoxylate	POCl ₃	7-Hydroxy-6-acetyl-trans-octalino- (2',3',4,3)-coumarin	_	96
ω-Chloro- resaceto-	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-6-chloroaceto- coumarin	9	126
phenone	Ethyl acetoacetate	HCl	7-Hydrnxy-4-methyl-6-chloroaceto- coumarin	4	126
	Diethyl oxalacetate	$Z_{D}Cl_{2} + HC$	1 7-Hydroxy-4-carhethoxy-6-chloro- acetocoumarin	45	126
2-Propionyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-propionyl- coumarin	_	204
4-Propionyl- resorcinol	Ethyl acetoacetate	POCl ₃	7-Hydroxy-4-methyl-6-propionyl- coumarin	25	12
	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-6-propionyl- coumarin	24	114
2-Butyryl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-hutyryl- coumarin	_	205
4-Butyryl- resorcinol	Ethyl acetoacetate	POCl ₃	7-Hydroxy-4-methyl-6-hutyryl- coumarin	30	12
	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-6-hutyryl- coumarin	37	114
4-Isovaleryl- resorcinol	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-6-isovaleryl- cnumarin	45	115
4-Lauroyl- resorcinol	Ethyl acetoacetate	AlCl3	5-Hydroxy-4-methyl-6-lauroyl- coumarin	27	115
4-Palmitoyl- resorcinol	Ethyl acetoacetate	AlCl3	5-Hydroxy-4-methyl-6-palmitoyl- coumarin	84	115
4-Stearoyl- resorcinol	Ethyl acetoacetate	AICl ₃	5-Hydroxy-4-methyl-6-stearnyl- coumarin	33	196
2-Benzoyl- resorciool	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-henzoyl- coumario	_	54
4.0	Diethyl acetosuccioate	POCI ₃	Ethyl 7-hydroxy-4-methyl- 8-benzoylcoumario-3-acetate		42
4-Benzoyl- resorciool	Ethyl acetoacetate	POCI3	7-Hydrnxy-4-methyl-6-beozoyl- coumarin	10	12
	Ethyl acetoacetate	AlCl ₃	5-Hydrnxy-4-methyl-6-beozoyl- cnumario	_	17
2-o-Toluyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-o-toluyl- coumarin		205
2-p-Toluyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4-methyl-8-p-toluyl- coumarin		204
4-p-Toluyl- resorcinol	Ethyl acetoacetate	AICI3	5-Hydroxy-4-methyl-6-p-toluyl- coumarin		114
4-Phenyl- acetyl- resorciool	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-6-phenyl- scetylcoumarin	42	114
4-Chloro- 6-methyl-	Ethyl acetoacetate	H ₂ SO ₄ ; P ₂ O ₅	5-Hydroxy-6-chloro-4,7-dimethyl- coumarin	_	43
resorciool	Ethyl α-methylacetoacetate	H ₂ SO ₄ ; P ₂ O ₅	5-Hydroxy-6-chloro-3,4,7-tri- methylcoumarin		43

Condensations with Dihydric Phenols

				Yield	Refer-
		Condensing	Product	%	ence
Phenol 4-Chloro-	Acid or Ester Ethyl a-ethylacetoacetate	Agent H ₂ SO ₄ ; P ₂ O ₅	5-Hydraxy-6-chloro-3-ethyl-4,7-di-	_	43
5-methyl-	Citric acid	H ₂ SO ₄	methylcoumarin 5-Hydroxy-6-chloro-7-methylcou-	_	43
(Cont'd)	Ethyl acetoacetate	H ₂ SO ₄ ; P ₂ O ₅	marin-4-acetic acid 5-Hydroxy-4-methyl-6(or 8)-chloro-	_	185
6-ethyl-			8-(or 6)-ethylcnumarin 5-Hydroxy-3,4-dimethyl-6(or 8)-		185
resorcinol	Ethyl α-methylacetoacetate	H ₂ SO ₄	chloro-8(or 6)-ethylcoumarin		43
4-Bromo- 5-methyl-	Ethyl acetoacetate	H ₂ SO ₄ ; P ₂ O ₅	5-Hydrnxy-6-bromo-4,7-dimethyl- cnumarin		43
resorcinol	Ethyl α-methylacetoacetate	H ₂ SO ₄	5-Hydrnxy-6-bromo-3,4,7-tri- methylcoumarin	_	
4-Chloro- 6-propionyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	5-Hydroxy-4-methyl-6(or 8)-chloro- 8(ar 6)-propionylcoumnrin	-	185
6-Bromn- 4-acetyl- resorcinol	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-6-acetyl- 8-bromocoumarin	16	117
	- Ethyl acetoacetate	H_2SO_4	5-Hydroxy-4,6,8-trimethylcoumarin	-	206
2-Methyl- 4-ethyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄ (80%)	o) 7-Hydroxy-4,8-dimethyl-6-cthyl- coumarin	-	22
2-Methyl- 4-propyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄ (80%	6) 7-Hydroxy-4,8-dimethyl-6-propyl- coumarin	-	23
2-Ethyl- 4-methyl- resorcinol		H ₂ SO ₄	7-Hydroxy-4,6-dimethyl-8-ethyl- coumarin	90	180
2-Ethyl- 5 methyl-	Ethyl acetoacetate	H ₂ SO ₄ (739	%) 7-Hydroxy-4,5-dimethyl-8-ethyl- coumarin	70	207
resorcino	Ethyl α-methylacetoacetat	e H ₂ SO ₄ (73		_	207
	Ethyl α-ethylaceto acetate	H ₂ SO ₄ (73		-	207
	Ethyl α-propylacetoaceta	te H ₂ SO ₄ (73	%) 7-Hydroxy-3-propyl-4,5-dimethyl-	-	207
2,4-Diethy resorcin		H ₂ SO ₄ ; C ₂ H ₅ O ₁	8-ethylcoumarin 7-Hydroxy-4-methyl-6,8-diethyl- Na coumarin	_	208
4-Ethyl- 5-methy	·-	H ₂ SO ₄ (8)	5%) 7-Hydrnxy-5-methyl-6-ethyl-	50	209
resorcit		H ₂ SO ₄ (8		60	209
4,6-Dieth resorci	nol	H ₂ SO ₄ (7	coumarin 5%) 5-Hydroxy-6,8-diethylcoumarin	_	210
	Ethyl acetoacetate	H ₂ SO ₄ (7	75%) 5-Hydroxy-4-methyl-6,8-diethyl- coumarin	_	. 210
	Ethyl cyclopentanone-2- boxylate		5-Hydroxy-6,8-diethylcyclopenten (1',2',4,3)-coumarin	10- 3	91
2-Propy	Ethyl 4-methylcyclopen none-2-carboxylate	ta- POCl3	5-Hydroxy-6,8-diethyl-4'-methyl- cyclopenteno-(1',2',4,3)-coumar		91
5-met	hyl-	H ₂ SO ₄ (73%) 7-Hydroxy-4,5-dimethyl-8-propyl		_ 207
	The thy tace to acc	,	(73%) 7-Hydrnxy-3,4,5-trimethyl- 8-propylcoumarin	-	_ 207
******	References 142-244 are listed m	n pp. 57-58.	- F-oppositioning		

Condensations with Dihydric Phenols

Phenol	Acid or Ester	Condensing Agent	Product	Yie %	
2-Propyl- 5-methyl-	Ethyl α -ethylacetoacetate	H ₂ SO ₄ (73%)		70 	
resorcinol (Cont'd)	Ethyl α-propylacetoacetate	H ₂ SO ₄ (73%)	7-Hydroxy-3,8-dipropyl-4,5-di- methylcoumarin	-	207
2,4-Dihy- droxy-	Ethyl acetoacetate	H ₂ SO ₄	5-Hydroxy-4-methyl-8-ethylcou- marin-6-carhoxylic acid	15	211
5-ethyl- benzoic acid	Ethyl acetoacetate	AlCl ₃	5-Hydroxy-4-methyl-8-ethylcou- marin-6-carhoxylic acid	24	211
Methyl 2,4-di- hydroxy-	Ethyl acetoacetate	H ₂ SO ₄ (73%)	Methyl 5-hydroxy-4-methyl- 8-ethylcoumarin-6-carboxylate	38	12
5-ethyl- benzoate	Ethyl acetoacetate	H ₂ SO ₄ (80%)	Methyl 5-hydroxy-4-methyl- 8-ethylcoumarin-6-carboxylete	22	211
	Ethyl acetoacetate	AlCl ₃	Methyl 5-hydroxy-4-methyl- 8-ethylcoumarin-6-carbo-ules	49	211
5-Methyl- resorcinol- 2-carhoxylic acid (p-or-	Ethyl acetoacetate	H ₂ SO ₄	7-Hydroxy-4,5-dimethylcoumarin- 8-carboxylic acid	32	212
sellinicacid) Ethyl	Malic acid	H ₂ SO ₄	5-Hudrom 7 - 43		
5-methyl-	Ethyl acetoacetate	H ₂ SO ₄	5-Hydroxy-7-methylcoumarin † Ethyl 5-hydroxy-4,7-dimethyl-	67	213
resorcinol-	Zinyi accidacetate		coumarin-6-carboxylata	60	213
6-carhox- ylate	Ethyl acetoacetate	AlCl ₃	Ethyl 5-hydroxy-4,7-dimethyl- coumarin-6-carboxylate	30	213
2,4-Dihy- droxy- 3-isoamyl- henzoic acid	Malic acid	H ₂ SO ₄	7-Hydroxy-8-isoamylcoumarin- 6-carboxylic acid	41	189
5-Methyl- 2-acetyl- resorcinol (γ-orca- ccto-	Ethyl acetoacctate	H ₂ SO ₄ ; H ₂ SO ₄ (73%); POCl ₃	5-Hydroxy-4,7-dimethylcoumarin ‡		214
phenone) 5-Methyl-	77/2 1	H ₂ SO ₄	5-Hydroxy-4,7-dimethylcoumarin ‡		
6-acetyl-	Ethyl acctoacetate Ethyl acctoacetate	POCl ₃	coumarin	18	17 12
(β-orcaceto- phenone)	Ethyl acctoacetate	AlCl ₃	5-Hydroxy-4,7-dimethyl-6-acetyl- coumarin	_	17
5-Methyl-2- propionyl- resorcinol	Ethyl acetoacetate	H ₂ SO ₄	5-Hydroxy-4,7-dimethylcoumarin † 5-Hydroxy-4,7-dimethylcoumarin §	_	215
5-Methyl- 2-hutyryl- resorcinol	Ethyl acetoacetate		5-Hydroxy-4,7-dimethylcoumarin f	-	215
2-Ethyl- 4-acetyl- resorcinol	Ethyl acetoacetate	POCl ₃	7-Hydroxy-4-methyl-6-acetyl- 8-ethylcoumarin	24	184

[†] A carbethoxyl group was eliminated in the condensation.

An acetyl group was eliminated in the condensation.

[§] A propionyl group was eliminated in the condensation.

A butyryl group was eliminated in the condensation.

Condensations with Dihydric Phenols

		CONDENSATION	S	VITH D.	LILL	MIC THERESE			
			Con	densing				-	Refer-
Phenol		Acid or Ester	-	gent		Product	9	%	ence
4-Ethyl-	Ethyl					lroxy-3,4-dimethyl-6-ethyl- cetylcoumarin		75	55
2-acetyl- resorcinol	Ethyl	a-ethylacetoacetate	H ₂ S	04 (73%)	7-Hy	droxy-3,6-diethyl-4-methyl- cetylcoumarin		70	55
	Ethyl	α-propylacetoacetate	H ₂ S	0 (73%)	7-Hy	droxy-3-propyl-4-methyl- thyl-8-acetylcoumarin		70	55
	Ethy	i α-hutylacetoacetate	H ₂ S	O ₄ (73%)	7-Hy	droxy-3-butyl-4-methyl- ethyl-8-acetylcoumarin			55
	Ethy	l α -allylacetoacetate	H25	304 (73%)	7-H3	droxy-3-allyl-4-methyl- ethyl-8-acetylcoumarin		50	55
	Ethy	l benzoylacetate	H_2	304 (73%)	7-H	ydroxy-4-phenyl-6-ethyl- acetylcoumarin		80	55
4-Ethyl- 6-acetyl-	Eth	l acetoácetate	PO	Cl ₂	5-H	ydroxy-4-methyl-6-acetyl- -ethylcoumarin			12
resorcinol	Eth	yl acetoacetate	AJ	Cl ₂	5-H	ydroxy-4-methyl-6-acetyl- -ethylcoumarin		39	117
4-Ethyl- 2-benzoyl-		yl acetoacetate	H	2SO4	7-F	iydroxy-4-methyl-6-ethyl- -benzoylcoumarin			216
resorcinol		nyl acetoacetate	H	2804 (73%)	7-I	Iydroxy-4-methyl-6-ethyl- 3-benzoylcoumarin		66	207
2,4-Diethyl- 5-methyl- resorcinol		hyl acetoacetate	A	IC13	7-1	Hydroxy-4,5-dimethyl-6,8-di- ethylcoumarin	•	-	207
4,6-Diethyl- 5-methyl	- M	alic acid	F	I ₂ SO ₄ (85%	5) 5-	Hydroxy-6,8-diethyl-7-methy coumarin	· <u>l</u> -	-	209
resorcino		thyl acetoacetate	3	H ₂ SO ₄ (85%	6) 5-	Hydroxy-4,7-dimethyl-6,8-di ethylcoumarin	-	-	209
Hydroquin	I	falic acid Ethyl acetoacetate Ethyl α-methylacetoacetat Ethyl α-methylacetoacetat	e	H ₂ SO ₄ H ₂ SO ₄ H ₂ SO ₄ P ₂ O ₆	6 6	Hydroxycoumarin -Hydroxy-4-methylcoumarin -Hydroxy-3,4-dimethylcoum:		Poor 20-34 3 30	39 148, 217 108, 217 4
		Ethyl a-ethylacetoacetate		AlCl ₃		-Hydroxy-2,3-dimethylchrom -Hydroxy-3-ethyl-4-methylco marin			207
		Diethyl acetonedicarhoxyl Diethyl oxnlacetate	ate	H ₂ SO ₄ H ₂ SO ₄		Ethyl 6-hydroxycoumarin-4-a Ethyl 6-hydroxycoumarin-4-o boxylate		Poor	89
		Ethyl cyclohexanone-2-car boxylate	•	H ₂ SO ₄		6-Hydroxy-3,4-cyclohexenoco marin	u-	10	9
		Ethyl 1-methylcyclohexas 3-one-4-carboxylate	1-	H ₂ SO ₄		6-Hydroxy-5'-methyl-3,4-cyc hexenocoumarin	lo-	2	
Hydro- quino diacet		Ethyl acetoacetate		H ₂ SO ₄ (7	3%)	6-Hydroxy-4-methylcoumari	n	30	
Hydro- quind mond meth ether	yl	Ethyl & methylacetoacet	ate	H ₂ SO ₄		6-Methoxy-3,4-dimethylcour	marin	Poo	r 218
	luinone	Ethyl acetoacetate		H ₂ SO ₄ (73%)	6-Hydroxy-4-methyl-7-chlor marin	-rocou	2	0 56
	hylhy-	Malic acid		H2SO4 (85%)	6-Hydroxy-7-methylcoums	rin	4	5 219
drox	quinone	Ethyl acetoacetate Ethyl c-methylacetoac	tate	H-SO.	(73%)	6-Hydroxy-4,7-dimethylcot	ımarin		0 56 5 56
		Ethyl α-ethylacetoacet	ate	H ₂ SO ₄		marin 6-Hydroxy-3-ethyl-4,7-dim		2	25 56
No	te: Refe	rences 142-244 are listed o	n pp.	57-58.		coumarin			

· TABLE II—Continued
Condensations with Dihydric Phenols

- %	спсе
	CHCo
,7-dimethyl- 20	56
-methyl- 45	56
-ethylcou- 45	56
yl-7-ethyl- 40	56
4-methyl- 35	56
methyl- 5-10	56
ethylcou- 15	56
arin	36
methyl- 17	220, 221
ntamethyl- 19	221
	methyl- 45 ethylcou- 45 d-rethyl- 40 4-methyl- 35 methyl- 5-10 ethylcou- 15 amyl- arin methyl- 17

Note: References 142-244 are listed on pp. 57-58.

TABLE III

CONDENSATIONS WITH TRIHYDRIC PHENOLS

Phenol	Acid or Ester	Condensing Agent	Product	Yiold %	Refer-
Pyrogallol	Malic acid Ethyl acetoacetate Ethyl acetoacetate Ethyl acetoacetate Ethyl acetoacetate Ethyl acetoacetate Ethyl acetoacetate Ethyl a-chloroacetoacetate	H ₂ SO ₄ H ₂ SO ₄ P ₂ O ₅ H ₃ PO ₄ FeCl ₃ ; TiCl ₄ SnCl ₄ H ₂ SO ₄ P ₂ O ₅	7.8-Dihydroxy-d-methylcoumarin 7.8-Dihydroxy-d-methylcoumarin 7.8-Dihydroxy-d-methylcoumarin 7.8-Dihydroxy-d-methylcoumarin 7.8-Dihydroxy-d-methylcoumarin 7.8-Dihydroxy-d-methylcoumarin 7.8-Dihydroxy-d-methyl- coumarin 7.8-Dihydroxy-d-chloro-d-methyl-	Quant.	1 2 33, 107 127 128 128 32
	Ethyl α-methylacetoacetate	H ₂ SO ₄ P ₂ O ₅ H ₂ SO ₄	coumarin 7,8-Dihydroxy-3,4-dimethylcou- marin	31	107
	Ethyl α -methylacetoacetate Ethyl α -ethylacetoacetate		7,8-Dihydroxy-3,4-dimethylcou- marin 7,8-Dihydroxy-3-ethyl-1-methyl-	Poor	33, 107 33
	Ethyl α-allylacetoacetate	POCl ₃	coumarin 7,8-Dihydroxy-3-allyl-4-methyl- coumarin	03	70
	Ethyl α-allylacetoacetate	HCI	7,8-Dihydroxy-3-(β-chloropropyl)- 4-methylcoumaria	47	124
	Ethyl α -(α -hydroxy- β , β , β - trichlorothyl)acetoacetate Ethyl α -(α -hydroxy- β , β , β - trichloroethyl)acetoacetato	H ₂ SO ₄ (78%) POCl ₃	7,8-Dihydroxy-3-(α-hydroxy-β,β,β- trichloroethyl)-4-methylcounarin 7,8-Dihydroxy-3-(α-hydroxy-β,β,β- trichloroethyl)-4-methylcoumarin	26 Quant	72 72

Condensations with Trihydric Phenols

	Condensation	s with T	RIHYDI	SIC PHENODS			
					Yield	Refe	
	/ <u>-</u>	Condensing		Product	%	enc	
Phenol	Acid or Ester	Agent	- TT - 3	xy-8-methoxy-4-methyl-	_	22	6
2,4-Dihy-	Ethyl acctoacetate	H_2SO_4					
droxyanisole		00 (000)	coum	rdroxycoumarin	30	22	
	mane acea	H ₂ SO ₄ (97%)	6,7-Din	droxy-4-methylcoumarin	92	60,	219
droquinone	Ethyl acetoacetate	H ₂ SO ₄	0,1-Din	All Orhad-Meanly 100 are			
triacetate		(73-					
		75%) ZnCl ₂	Ethyl f	,7-dihydroxycoumarin-	_	1	24
	Diethyl oxalacetate	ZnCi2	A.cat	hoxylate			
	mu 11-1	H ₂ SO ₄ (80%)	67-Dil	ydroxy-3-phenylcoumarin		2	28
	Ethyl hydroxymethylene phenylacetate	112501 (5070)					29
7011t-al	Ethyl acetoacetate	H ₂ SO ₄	5.7-Di	hydroxy-4-methylcoumarin			:29 170
Phiotoginginor	Ethyl acetoacetate (3 moles)	H ₂ SO ₄	Trime	thyltricoumarin	10	,	101
	Ethyl acetoacetate	P2O5	5.7-Di	hydroxy-4-methylcoumarin			127
	Ethyl acetoacetate	H ₃ PO ₄	5,7-D	hydroxy-4-methylcoumarir	, -		128
	Ethyl acetoacetate	FeCl ₃	5,7-D	ihydroxy-4-methylcoumarii	n 0		128
	Ethyl acetoacetate	SnCl4	5.7-D	ihydroxy-4-methylcoumarii	n 1		26
	Ethyl a-chloroacetoacetate	H_2SO_4	5,7-D	ihydroxy-3-chloro-4-methy	1- 3	7	20
				marin			33
	Ethyl a-chloroacetoacetate	P_2O_5		hydroxy-3-chloro-4-methy	1-	-	00
			CO	ımarin	_	_	101
	Ethyl α-methylacetoacetate			Dihydroxy-3,4-dimethylcou-	_		
		(75%);	m	arin			
		P ₂ O ₅		nn 1 n 11-14-48-6	1	46	101
	Ethyl α -ethylacetoacetate	H ₂ SO ₄		Dihydroxy-3-ethyl-4-methy	1-	•••	
		(73%);	C	oumarin			
	Ethyl a-allylacetoscetate	P ₂ O ₅ H ₂ SO ₄	57.	Dihydroxy-3-allyl-4-methyl	-	72	70
	Ethyl a-shylacetoscetate	112504		oumarin			
	Ethyl a-allylacetoacetate	HCl		-Dihydroxy-4-methyl-3-(β-0	hloro-	_	124
	11000 1 00 0000 1	201		propyl)coumarin			
	Ethyl α-(α-hydroxy-β,β,β	3- P ₂ O ₅		-Dihydroxy-3-(α-hydroxy-f	3,β,β-	Poor	72
	trichloroethyl)acetoace			trichloroethyl)-4-methylcou			#0
	Ethyl α-(α-hydroxy-β,β,	8- POCl3		7-Dihydroxy-3-(α-hydroxy-		29	72
	trichloroethyl)acetoace	etate		trichloroethyl)-4-methylcou	marin		105
	Ethyl a-phenylacetoacet	ate ZnCl ₂	5,	7-Dihydroxy-3-phenyl-4-me	thyl-	_	100
	7 .1			coumarin			105
	Ethyl a-benrylacetoacet	ate H ₂ SO ₄	5	7-Dihydroxy-3-henzyl-4-me	thyl-	_	100
	Ethyl a-benzylacetoace	tate POCla		coumarin	. + b l .		172
	23.137 G-06.123 (266.030)	tate FOCIŞ	0	,7-Dihydroxy-3-benzyl-4-me coumarin	thyl-		
	Ethyl a-o-carboxybenz	yl- HCl	,	5,7-Dihydroxy-3-o-carboxyb	enzvl-	Good	79
	acetoscetate	,		4-methylcoumarin			
	Diethyl acetorucciuste	H ₂ SO ₄		Ethyl 5,7-dihydroxy-4-meth	vl-	_	179
				coumarin-3-acetate			
	Diethyl acetosuccinate		(80%)	5.7-Dihydroxy-4-methylcou	marin	_	34
	Diethyl acetosuccinate	POC1	3	Ethyl 5.7-dihydroxy-4-meth	ıyl-	91	34
	Diabet			coumarin-3-acetate			***
	Diethyl o-acetylgiuta		•	5.7-Dihydroxy-4-methylcou	ımarin-	32	77
		•	n∝d. and	3-propionic acid			
	Ethyl phthalylaceton	789 cetate HCl	(a)	57 Dh			79
	, ,	1101		5,7-Dihydroxy-4-methylco		_	• •
	Acrtonedicarboxylie	acid H ₂ S0	D ₄	3-benzoyl-o-carboxylic a 5,7-Dihydroxycoumarin-4-		_	26
			•	acid	act with		
	Ethyl butyroscetate		0. (75%)	5,7-Dihydroxy-4-propylco	umarin	_	35
N	ete: Pelereures 142-244 are listed	on to. 57-59					
		- , ,					

THE PECHMANN REACTION

TABLE III-Continued

CONDENSATIONS WITH TRIHYDRIC PHENOLS

		Condensin	σ.	Yield	i Refer
Phenol	Acid or Ester	Agent	Froduct	7 TIEIG	neier ence
Phloroglucino	Ethyl benzoylacetate	P ₂ O ₅	5,7-Dibydroxy-4-pbenylcoumarin	~·	101
(Cont'd)	Ethyl benzoylacetate	ZnCl ₂	5,7-Dihydroxy-4-phenylcoumarin		222
	Ethyl a-benzylhenzoylacetat		5,7-Dihydroxy-3-henzyl-4-phenyl- coumarin	85-90	
	Ethyl 3,4,5-trimethoxy- benzoylacetate	H ₂ SO ₄ (73%	5) 5,7-Dihydroxy-4-(3',4',5'-trimetb- oxypbenyl)coumarin	_	223
	Etbyl γ-phenylacetoacetate	H ₂ SO ₄ (concd. and 80%)	5,7-Dibydroxy-4-benzylcoumarin	_	80, 81
	Etbyl cyclopentanone-2-car- boxylate	POCl ₃	5,7-Dihy droxy cy clopenteno- (1',2',4,3)-coumarin	55	91
	Etbyl 4-metbylcyclopenta- none-2-carboxylate	POCl ₃	5,7-Dibydroxy-4'-methylcyclo- penteno-(1',2',4,3)-coumarin	_	91
	Etbyl cyclohexanone-2-car- boxylate	POCl ₃	5,7-Diby droxycyclohexeno- (1',2',4,3)-coumarin	_	124
	Ethyl 4-metbylcyclohexa- none-2-carboxylate	POCl ₃	5,7-Dibydroxy-4'-methylcyclo- bexeno-(1',2',4,3)-coumarin	-	97
	Etbyl 5-metbylcyclohexa- none-2-carboxylate	H ₂ SO ₄ ; POCl ₃	5,7-Dihydroxy-5'-metbylcyclo- hexeno-(1',2',4,3)-coumarin	_	97
	Etbyl 5-metbylcyclohexa- none-2-carhoxylate	ZnCl ₂	3,4-Tetrabydro-4'-met bylbenzo- 5,7-dihydroxycoumarin	75-80	94
	Etbyl 6-metbylcyclohexa- none-2-carboxylate	POCl ₃	5.7-Dibydroxy-6'-metbylcyclo- hexeno-(1',2',4,3)-coumarin	_	97
	Ethyl trans-β-decalone- 3-carboxylate	H ₂ SO ₄	5,7-Dihydroxy-trans-octalino- (2',3',4,3)-coumarin	_	97
	Ethyl \$\beta\$-coumaranone-2-car- boxylate	HCI	5,7-Dihydroxycoumarono-(2',3',3,4)- coumarin	_	100
	Ethyl 5-metbyl-β-coumara- none-2-carboxylate	HCl	5,7-Dibydroxy-5'-metbylcoumarono- (2',3',3,4)-coumarin	_	100
	Ethyl 6-metboxy-β-coumara- none-2-carboxylate	HCI	5,7-Dihydroxy-6'-methoxycouma- rono-(2',3',3,4)-coumarin	_	100
	Ethyl 3-bydroxy-7-methoxy- 3-chromene-4-carboxylate	HCI	5,7-Dihydroxy-7'-methoxychro- meno-(3',4',4,3)-coumarin		99
	Etbyl 3-bydroxy-8-metboxy- 3-chromene-4-carboxylate	H ₂ SO ₄ (85%); HCl	5,7-Dihydroxy-8'-methoxychro- meno-(3',4',4,3)-coumarin (impure)	_	99
	Ethyl 3-hydroxy-6,7-dimeth- oxy-3-chromene-4-car-	H ₂ SO ₄ (85%)	5,7-Dihydroxy-6',7'-dimethoxy- chromeno-(3',4',4,3)-coumarin (impure)	_	99
Phloroglucinol mono- methyl ether	boxylate Ethyl acetoacetate	H ₃ PO ₄	5-Hydroxy-7-methoxy-1-methyl- coumarin and 7-bydroxy-5-meth- oxy-1-methylcoumarin	_	230
Phloroglucinol	Ethyl acetoacetate	P ₂ O ₅	5,7-Dimethoxy-4-methylcoumarin	70	101
dimethyl	Ethyl acetoacetate	H ₃ PO ₄	5,7-Dimethoxy-1-methylcoumarin	63	230
ether	Ethyl α-methylacetoacetate	P ₂ O ₅	5,7-Dimethoxy-3,4-dimethylcou- marin	-	101
	Ethyl α-benzylacetoacetate	P ₂ O ₅	5,7-Dimethoxy-3-benzyl-4-methyl- coumarin		231
	acetoacetate	P ₂ O ₅	5,7,4'-Trimethoxy-3-benzyl-4-metb- ylcoumarin		231
Methyl- phloro- glucinol	Malic acid	H ₂ SO ₄	Isolated as 5,7-dimethoxy-8-methyl- coumarin and 5,7-dimethoxy- 6-methylcoumarin after methyla- tion	11 :	232

Condensations with Trihydric Phenols

	CONDENSAI	Condensing		Yield %	Refer- ence
Phenol	Acid or Ester	Agent H ₂ SO ₄	Product 5,7-Dihydroxy-1,6-(or 8)-dimethyl-	95	58
phloro-	Ethyl acetoacetate	112004	coumarin		
glucinol (Cont'd) Dimethyl-	Ethyl acetoacetate	H ₂ SO ₄	5,7-Dihydroxy-4,6,8-trimethyl- coumarin	69	58
phloro- glucinol Methyl	Ethyl acetoacetate	H ₂ SO ₄ (80%)	Methyl 5,7-dihydroxy-4-methyl-	47	59
phloro-	•	AlCla	coumarin-6(or 8)-carboxylate Methyl 5,7-dihydroxy-4-methyl-	44	59
glucinol carboxylate	Ethyl acetoacetate		coumarin-6(or 8)-carboxylate 5,7-Dihydroxy-4-methyl-6(or 8)-	18	17
Phloroaceto- phenone	Ethyl acetoacetate	H ₂ SO ₄	acetylcoumarin	18	17
риенопо	Ethyl acetoacetate	AlC1 ₃	5,7-Dihydroxy-4-methyl-6(or 8)- acetylcoumarin	,,,	207
Phlorobenzo- phenone	Ethyl acetoacetate	H ₂ SO ₄ (85%		-	201
	140 Ott Usted or	nn 57_59			

Note: References 142-244 are listed on pp. 57-58.

TABLE IV

Condensations with Naphthols *

	CONDENSA	.10110 11111			
Phenol	Acid or Ester	Condensing Agent	Product	Yield %	Refer- ence 233
α-Naphthol	Ethyl acetoacetate Ethyl acetoacetate	H ₂ SO ₄ H ₂ SO ₄ H ₂ SO ₄ (80-84%)	α-Naphthacoumarin 4-Methyl-α-naphthacoumarin 4-Methyl-1,2,α-naphthapyrone	60 85- Quant. 18	233, 234 108, 156 33, 108
	Ethyl acetoacetate Ethyl acetoacetate Ethyl acetoacetate	P ₂ O ₆ HCl H ₃ PO ₄ ;	4-Methyl-1,2,α-naphthapyrone 4-Methyl-1,2,α-naphthapyrone 4-Methyl-1,2,α-naphthapyrone	93	123, 234 127
	Ethyl α -chloroacetoacetate	CH ₃ CO ₂ N ₃ H ₂ SO ₄	3-Chloro-4-methyl-1,2,α-naphtha- pyrone	Good	26, 159
	Ethyl α -chloroacetoacetate Ethyl α -methylacetoacetate	P ₂ O ₅ H ₂ SO ₄	3-Chloro-4-methyl-1,2,α-naphtha- pyrone 3,4-Dimethyl-α-naphthacoumarin	_	33 33, 75,
	Ethyl a-methylacetoacetate Ethyl a-ethylacetoacetate	(concd. or 84%)	3,4-Dimethyl-c-naphthacoumarin 3-Ethyl-4-methyl-c-naphthacou-	33 30	108 33, 108 233
	Ethyl α-propylacetoacetate		marin 3-Propyl-4-methyl-1,2,α-naphtha- pyrone	_	33
	Ethyl α-isopropylaceto- acetate	H ₂ SO ₄	3-Isopropyl-4-methyl-1,2, α -naph- thapyrone	_	33

Note: References 142-244 are listed on pp. 57-58.

• The coumarins and chromones derived from naphthols have been called α - or β -naphthacoumarins or α - or β - or chromones by various workers. These names are inappropriate as they do not convey the proper idea of the structures of these compounds. The names 1,2,α-naphthapyrone and 1,4,α-naphthapyrone for the coumarins and chromones. respectively, from α -naphthol and 1,2, β , α -naphthapyrone and 1,2, β , β -naphthapyrone for the coumarins from β -naphthol, and 1,4,6, α -naphthapyrone for the coumarins from β -naphthol. and $1.4.\beta.\alpha$ -naphthapyrone and $1.4.\beta.\beta$ -naphthapyrone for the chromones from β -naphthapyrone and $1.4.\beta.\beta$ -naphthapyrone for the chromones from β -naphthal as suggested by Dey and Lakshminarayan (ref. 110) are rational. However, in order to avoid confusion, the original names as given by the authors are given in the tables.

TABLE IV—Continued

Condensations with Naphthols

Phenol	Acid or Ester	Condensing Agent	; Product	Yield %	Refer-
α-Naphthol (Cont'd)	Ethyl α -allylacetoacetate	H ₂ SO ₄	3-Allyl-4-methyl-5,6-naphtha- α-pyrone	86	70
	Ethyl α -allylacetoacetate	HCl	3-β-Chloropropyl-4-methyl- 5,6,α-naphtha-1,2-pyrone	_	124
	Ethyl α-(α-hydroxy-β,β,β- trichloroethyl)acetoace- tate	POCl ₃	4-Methyl-3-(α-hydroxy-β,β,β-tri- chloroethyl)-1,2,α-naphthapy- rone	25	72
	Ethyl α-phenylacetoacetate	H ₂ SO ₄	3-Phenyl-4-methyl-1,2,α-naphtha pyrone †		104
	Ethyl α-benzylacetoacetate	H_2SO_4	3-Benzyl-4-methyl-1,2,α-naphtha- pyrone †	-	102
	Ethyl α -benzylacetoacetate	POCl ₃	3-Benzyl-4-methyl-1,2,α-naphtha- pyrooe	_	172
	Diethyl acetosuccinate	H_2SO_4	Ethyl 4-methyl-1,2,α-naphtha- pyrone-3-acetate	24	34, 75, 76
	Diethyl acetosuccinate	H ₂ SO ₄ (80%)	111	_	34
	Diethyl acetosuccinate	P_2O_5	Ethyl 4-methyl-1,2,α-naphtha- pyrone-3-acetate	-	34
	Diethyl acetosuccinate	POCl ₃	4-Methyl-1,2,α-naphthapyrooe- 3-acetic acid	40	34
Diethyl α-acetylgl	Diethyl acetosuccinate	AlCl ₃	4-Methyl-1,2,α-oaphthapyrone- 3-acetic acid	-	34
	Diethyl α -acetylglutarate	$\mathrm{H}_2\mathrm{SO}_4$	Ethyl 4-methyl-1,2,\alpha-naphtha- nyrone-3-propionate	27	77
	Diethyl α -acetylglutarate	H ₂ SO ₄ (78%)	4-Methyl-1,2,α-naphthapyrone-		77
	Ethyl γ-bromoacetoacetate	H_2SO_4	4-Bromomethyl-1,2,α-naphtha- pyrone	13	83
	Acetonedicarboxylic ocid	H ₂ SO ₄	1,2, a-Naphthapyrooe-4-acetic acid	Good	26
	Diethyl oxalacetate	H ₂ SO ₄	Ethyl α-naphthaeoumarin-4-car- boxylate		233 35
	Ethyl butyroacetate	H ₂ SO ₄ (75%)	α-Naphtha-4-propyl-α-pyrone 3-Benzyl-4-pheoyl-1,2,α-naphtha-	_	103
	Ethyl α-benzylbenzoyl- ncetate	H ₂ SO ₄ ; SnCl ₄	nvrone	_	81
	Ethyl 7-phenylacetoacetate	H ₂ SO ₄ (80%)	α-Naphtha-4-benzyl-α-pyrone Cyclopenteoo-(1',2',4,3)-1,2,α-	71	91
	Ethyl cyclopentanone-2-car- boxylate	H2SO4	naphthapyrone 4'-Methylcyclopeoteno-(1',2',4,3)-		91
	Ethyl 4-methylcyclopenta- none-2-carboxylate	H ₂ SO ₄	1,2.a-naphthapyrooe 3,4-Tetrahydrobeoronaphtha-	Quant.	94
	Ethyl cyclohexanone-2-car- boxylate	H ₂ SO ₄	coumarin 4'-Methylcyclohexeno-(1',2',4,3)-	Quant,	97
	Ethyl 4-methylcyclohexa- nooe-2-carboxylate	H:SO4	1.2.a-naphthapyrone 5'-Methylcyclohexeno-(1',2',4,3)-		97
	Ethyl 5-methylcyclohexa- none-2-carboxylate	11 ₂ SO ₄ ; POCl ₂	1,2,6-naphthapyrone 6'-Methyleyclohexeoo-(1',2',4,3)-	_	97
	Ethyl 6-methylcyclohexa-	POCl ₃	1,2,a-maphthapyrone	_	
	none-2-carboxylate Ethyl trans-β-decalone-	H\$204	frans-Octalino-(2',3',4,3)-1,2,a- naphthapyrone		97
-Chloro-	3-carboxylate		6-Chloro-1.2.a.8-naphthapyrone		61
a-naphthol	Malic acid Ethyl acetoacetate	11:504	6-Chloro-4-methyl-1,2,o.8-naph-	91	CI
maparation	Linja accioscente		thatyrone		

Note: References 142-244 are listed on pp. 57-55. The 1,4,a-naphthapyrone structure originally assigned to this compound is incorrect; refs. 165, 105.

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Condensations with Naphthols

		CONDENSA	TION	S WIT	нΝ	APHTHOLS			*
		•					Yiel	u -	efer-
				ensing		Product	%	•	ence
Phenol		Acid or Ester		ent	- 011	oro-4-methyl-1,2,α,β-naph			61
4-Chloro-	Ethyl a	cetoacetate	P205;			pyrone			
α -naphthol				I ₅ ONa;	Į.D:	ругоне			.1
(Cont'd)				CO2Na	2 6.1	Diehloro-4-methyl-1,2, α , β -	_	•	61
	Ethyl	α-chloroacetoacetate	H250	4; P ₂ O ₅	נייט,נט	phthapyrone		_	61
		O. Leading adults	ים.פר	١.	6-C)	aloro-3,4-dimethyl-1,2, α , β -	41	3	01
	Ethyl	α-methylacetoacetate	H ₂ SC	74	70	nhthanyrone			61
	To 1		P20		6-C	hloro-2,3-dimethyl-1,4, α , β -	-	-	U.
	Ethyl	α-methylacetoacetate	1 20	•	n	aphthapyrone			61
	T4hm	l α-ethylacetoacetate	H ₂ S	04	6-C	hloro-3-ethyl-4-methyl-	-	-	~-
	Eithy	t te-ethylacetolacetolac	11 20	••	1	.2. c. 6-naphthapyrone			61
	Fthu	l α-ethylacetoacetate	P20	a)	6-0	hloro-2-methyl-3-ethyl-	•	_	•-
	Lieny	La-cenj moreonomic		•		4.c.8-naphthapyrone			61
	Ethy	ylα-propylacetoacetate	H25	304	6-0	Chloro-3-propyl-4-methyl-			-
		, , ,				1.2. a. B-naphthapyrone		_	61
	Eth	yl α-propylacetoacetate	P ₂ (0 ₅	6-	Chloro-2-methyl-3-propyl-		_	•
						1,4,α,β-naphthapyrone			61
	Eth	ıylα-isohutylacetoaceta	te H2	\$0₄	6-	Chloro-3-isohutyl-4-methy	\ -		
						1,2,α,β-naphthapyrone	,	_	61
	Et	hylα-isohutylacetoacet:	te P	2Os	6	Chloro-2-methyl-3-isohuty	I -		
						1,4, \alpha, \beta-naphthapyrone			61
	Et	hyl α-phenylacetoaceta	te H	2SO4	6	-Chloro-3-phenyl-4-methyl			
	_					1,2,α,β-naphthapyrone	_	_	61
	E	thyl α-benzylacetoacets	te H	12SO4	'	Chloro-3-benzyl-4-methyl	-		
	***			T.00.		1,2,α,β-naphthapyrone Ethyl 6-chloro-4-methyl-1,	2.α.β-	_	61
	ш	lethyl acetosuccinate	1	H ₂ SO ₄		naphthapyrone-3-acetate			
	τ.	Diethyl acetosuccinate	,	H ₂ SO ₄		Ethyl 6-chloro-4-methyl-1,	2.α.β-		42
		olemyl acetosuccinate		Hånof		naphthapyrone-3-acetate	3		
						6-Chloro-4-methyl-1,2,α,β-	naph-	-	
						thapyrone-3-acetic acid			42
		Diethyl acetosuccinate		H2SO4 (8	(%0	6-Chloro-4-methyl-1,2,α,β	-naph-		74
						thapyrone-3-acetic acid			61
		Acetonedicarboxylic ac	id	$H_{2}SO_{4}$		6-Chloro-1,2,α,β-naphtha	yrone-	_	٧.
						4-acetie acid			61
		Ethyl benzoylacetate		H_2SO_4		6-Chloro-4-phenyl-1,2,a,f	l-naph-	_	
4-Bros		To 1		** **		thapyrone	a L		61
	no- aphthol	Ethyl acetoacetate		H ₂ SO ₄ ;	P2O5	6-Bromo-4-methyl-1,2,a,	3-napn-		
C-II.	арпиног	Ethyl α-methylacetos	notata	H ₂ SO ₄		thapyrone) ~ R-		61
		maji a-memjiacetoa	cetate	112504		6-Bromo-3,4-dimethyl-1, naphthapyrone	Δ,α,μ-		
		Ethyl α-methylaceto:	cetate	P2O6		6-Bromo-2,3-dimethyl-1,	4.α.β-		61
		•		-2-0		naphthapyrone	-1		
		Ethyl a-benrylaceto:	cetate	H ₂ SO ₄		6-Bromo-3-benzyl-4-met	hyl-	_	_ 61
						1,2,α,β-naphthapyron			12
	etyl-	Ethyl acetoacetate		H_2SO_4		4-Methyl-1,2,α-naphtha	pyrone ‡	-	_ 14
α-	-naphthol	Fibral		POC	n_3				_ 117
4.P	ropionyl-	Ethyl acetoacetate Ethyl acetoacetate		AIC13		4-Methyl-1,2,α-naphth:		-	_ 12
	-naphthol			H ₂ SO		4-Methyl-1,2, α -naphth:	apyrone §	_	-
_		Ethyl acetoacetate		PO AlCl ₃		436-43-10 c=34	- marona R	_	117
4-E	Butyryl-	Ethyl acetoacetate		POCI		4-Methyl-1,2,α-naphth			12
	z-caphtho			- 50	•	4-Methyl-1,2,α-naphth	apjione II		
	Note: Refe	erences 142-244 are liste	d on ~-	. K7 E0					

An acetyl group was eliminated in the condensation.

A propionyl group was eliminated in the condensation.

A butyryl group was eliminated in the condensation.

THE PECHMANN REACTION

TABLE IV-Continued

CONDENSATIONS WITH NAPHTHOLS

Phenol	4.23 - 70.5	Condensing		Yield	Refer
	Acid or Ester	Agent	Product	%	ence
β -Naphthol	Malic acid	H₂SO₄	β-Naphthacoumarin	Poor	39
	Ethyl acctoacctato	H ₂ SO ₄	4-Methyl-1,2,β,α-oaphthapyrone	20-39	234, 23
	Ethyl acctoacctate	112504	4-Mcthyl-1,2,β,α-naphthapyrone	25	110
			2-Methyl-1,4,β,α-naphthapyrone (isolated as styryl derivative)	Traces	
	Ethyl acetoacetate	H ₂ SO ₄ (80%)	4-Methyl-1,2,β,α-naphthapyrone	70	155
	Ethyl acctoacctate	P2O5	2-Methyl-1,4,β,α-naphthapyrone	10	110
	Ethyl acctoacctato	CH ₃ CO ₂ Na	4-Methyl-1,2,β,α-naphthapyrooc		127
			2-Methyl-1,4,β,α-naphthapyrone	-	
	Ethyl α-methylacetoacetate	H ₂ SO ₄	3,4-Dimethyl-1,2,β,α-naphtha- pyrone	_	71
	Ethyl a-methylacetoacetate	P2O5	2,3-Dimethyl-1,4,β,α-naphtha- pyronc	_	71
	Ethyl a-ethylacctoacetate	P_2O_5	2-Methyl-3-ethyl-1,4,β,α-naphtha- pyrone	_	71
	Ethyl α-propylacetoacetate	P_2O_5	2-Methyl-3-propyl-1,4,β,α-naph- thapyrone	_	71
	Ethyl α-isopropylaceto- acetate	P ₂ O ₆	2-Methyl-3-isopropyl-1,4,β,α- naphthapyτone	-	71
	Diethyl formylsuccinate	H ₂ SO ₄	8-Naphthapyrooe-3-acetic acid	_	76
	Diethyl acctosuccioate	H ₂ SO ₄	4-Methyl-β-naphthapyrooe- 3-acetic acid	40	34, 76
	Ethyl y-hromoacetoacctate	H ₂ SO ₄	4-Bromomethyl-β-caphthapyroce	_	83
	Acetonedicarhoxylic acid	H ₂ SO ₄	4,3,β-naphthapyτοoe-1-acetic acid	_	26
	Diethyl oxalacetate	H ₂ SO ₄	Ethyl 4,3,8-naphthapyrone- carhoxylate	_	26
	Ethyl beozoylacetate	P_2O_5	β-Naphthoffavooe	30	236
	Ethyl cyclopentaoooe- 2-carboxylate	P ₂ O ₅	Cyclopeoteno-(1',2',2,3)-1,4-β,α- naphthapyrooe	_	11
1,5-Dihydroxy- naphthaleos	Ethyl acctoacetate	HCl	6'-Hydroxy-4-methyl-7,8-henzo- coumarin	92	237
	Ethyl acetoacetate	AlCl ₃	6'-Hydroxy-4-methyl-7,8-henzo- coumarin	78	237
	Diethyl α-acetylglutarate	H ₂ SO ₄	Diethyl 4,4'-dimethylnaphtha- dipyrone-3,3'-dipropionate	_	930

Note: References 142-244 are listed on pp. 57-58.

TABLE V

Condensations with Miscellaneous Compounds

Compound	Acid or Ester	Con- densing Agent	Product	7514 %	Matro
1,2,3-Trihydroxy- 4-methoxybenzene	Malic acid	H ₂ SO ₄	6,7,8-Trihydroxycoumarin •	,	239
Lecanoric acid Thiopheool	Malic acid Methyl α-methylaceto- acetate	H ₂ SO ₄ P ₂ O ₅	5-Hydroxy-7-methylcoumarin 2,3-Dimethyl-1-thiochromom;	17	212 217
Thiotolenol	Ethyl acetoacetate	H_2SO_4	4,6-Dimethylthiopheny-1,2-pyrres	-	16

Note: References 142-244 are listed on pp. 57-58.
• Demethylation took place during the reaction.

TABLE V—Continued Condensations with Miscellaneous Compounds

00.					
		Con-		Yield	Refer-
Compound Ethyl 2-methyl-4-hy- droxythiophene- 3-carboxylate	Acid or Ester	densing Agent	Product	%	ence
		H ₂ SO ₄	Ethyl 4,6-dimethyl-5-thiocoumarin-	31	65
	Ethyl acetoacetate	_	7-carboxylate		
	Ethyl a-methylaceto-	H-SO4	Ethyl 3,4,6-trimethyl-5-thiocou-	_	65
	acetate		marin-7-carboxylate		65
	Diethyl acetylsuccinate	H ₂ SO ₄	Ethyl 4,6-dimethyl-5-thio-7-car-	_	•
			bethoxycoumarin-3-acetate	17	65
	Acetonedicarboxylic acid	H ₂ SO ₄	6-Methyl-7-carhethoxy-5-thiocou- marin-4-acetic acid		
	Ethyl o-cyclohexanone-	H ₂ SO ₄	Ethyl 3,4-cyclohexenyl-6-methyl-	46	65
	carboxylate	112004	5-thiocoumarin-7-carboxylate		63
7-Hydroxycoumarin	Malic acid	H ₂ SO ₄	Coumaro-7,6(or 7,8)-a-pyrone	60	62
	Malic acid	H2SO4	Coumarino-7,8, a-pyrone	53 3	U-2
			Coumarino-7,6, a-pyrone		62
7-Hydroxy-4-methyl-	Malic acid	H ₂ SO ₄	4-Methylcoumarino-7,8, a-pyrooe	70	63
coumarin	Malic acid	H_2SO_4	4-Methylcoumaro-7,6(or 7,8)-a-	•	
	Ethyl acetoacetate	H ₂ SO ₄	pyrone 4.4'-Dimethylcoumaro-7,6(or 7,8)-α-	30	63
	Emy accommente	112004	DALOUG DALOUG		
	Ethyl acetoacetate	H2SO4	4,4'-Dimethylcoumarino-7,8,a-	15	62
			pyrone	30	241
7-Hydroxy-3-chloro-	Malic acid	H ₂ SO ₄	3-Chloro-4-methylcoumaro-	30	212
4-methylcoumarin \$-Hydroxy-7-methyl-			7,6(7,8)-а-рутоое	50	63
coumarin	- Malie acid	H ₂ SO ₄	7-Methylcoumaro-5,6,α-pyrooe	•	
5-Hydroxy-4,7-dime	th- Malic acid	H ₂ SO ₄	4,7-Dimethylcoumaro-5,6,α-pyτooe	65	63
ylcozmarin	Ethyl acctoacetate	H ₂ SO ₄		15	63
*** *			pyrone		241
5-Hydroxy-3-chloro 4,7-dimethylcou-	 Malie acid 	H ₂ SO ₄	3-Chloro-4,7-dimethylcoumaro-5,6,0-	. 20	211
marin			pyrone		
7,5-Dihydroxycou-	Malic acid	H2SO4	8-Нудгохусоциаго-7,6,α-ругоос	40	63
marin		11500	8-113 th 013 (00 (11 th 10 th		
7.8-Dibydroxy-4-m	eth- Malic acid	H2SO.	8-Hydroxy-4-methylcoumaro-	55	63
7looumana 7,8-Dihydroxy-	14 12		7,6,α-рутопе		241
3-chloro-1-meth	Malie acid	H2SO			2
comaria	,		maro-7,6,α-pyrone		
5.7-Dibydrosy-	Malic zeid	H ₂ SO	5-Hydroxy-1-methylcoumaro-7,8,α-	60	63
4-methylocoma	ria		pyrone or 7-hydroxy-4-methyl-		
6'-Hydroxy-4-me	that the same		coumaro-5,6,0-pyrone		_ 237
7,8-benzooran	thyl- Ethyl acetoscetate	H ₂ SC			_ 231
2,2-Dimethyl-7-h	5- Malic acid		(%) coumarin		242
droxychroman	cte	H ₂ S(O4 Dimethyldihydroxypyronocoumarii	а –	
7(T)-History-2,		ZnC	l2 4.6.6.8-Tetramethyl-6.7-dihydro-	_	_ 121
trimethyl-3,4- dropulacime	y-		quinocoumarin		
6-Hid-myer	aran Malic add	** *			1 243
		H-2		n 5	1 243
£.7.Dirydroxye	 Malic acid 	H ₂ 9	(4',5-dihydropsoralen) (4',5'-Dihydro-8-hydroxy-2',3',7,6-	3	0 244
trans.			furocoumarin (4',5'-dibydro-		
Note: Data	ore 142-214 are leted on on		(lexotedance		
A COMMITTEE STATE AND ADDRESS OF THE PERSON	" " I la " . I I hip total on on				

Note: Paternoves 142-244 are listed on pp. 57-58.

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CHAPTER 2

THE SKRAUP SYNTHESIS OF QUINOLINES

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INTRODUCTION

Koenigs ¹ first synthesized quinoline in 1879 by passing allylaniline over heated litharge. Shortly thereafter ² he prepared quinoline by heating the condensation product of aniline and acrolein, thus antici-

¹ Koenigs, Ber., 12, 453 (1879).

² Koenigs, Ber., 13, 911 (1880).

pating the classical Skraup synthesis. This synthesis involves a series of reactions brought about by heating a primary aromatic amine, in which at least one position ortho to the amino group is unsubstituted, with glycerol, sulfuric acid, and an oxidizing agent. The product is a quinoline containing only those substituents that were originally present in the aromatic amine. Quinolines substituted in the hetero ring may be obtained by a modified Skraup synthesis in which a substituted acrolein or a vinyl ketone is used in place of glycerol.

MECHANISM

The Skraup reaction takes place through four successive steps: dehydration of glycerol to acrolein under the influence of sulfuric acid; addition of the aromatic amine to acrolein to form an intermediate β -arylaminoaldehyde (III); ring closure by dehydration to form 1,2-dihydroquinoline (IV); and oxidation of IV to quinoline (V). The re-

$$\begin{array}{c} CHO \\ CHO \\ CH_2 \\ CH_2$$

placement of glycerol by acrolein in the reaction with aniline, sulfuric acid, and an oxidizing agent under ordinary conditions results in much resinification and only a little quinoline.3 However, a high yield of quinoline can be obtained by passing acrolein vapor into the solution of aniline, sulfuric acid, and an oxidizing agent under proper conditions. 4, 5 The nitroanilines and the nitromethoxyanilines react readily with liquid acrolein to give good yields of the corresponding substituted quinolines, 6.7,8 especially when sulfuric acid is replaced by phosphoric acid.7

- ² Manske, unpublished observations.
- ⁴ Tchitchibabin, Swiss pat. 240,991 (1946).
- Kulka, unpublished observations.
- ⁶ Yale, J. Am. Chem. Soc., 69, 1230 (1947).
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- ⁸ Yale, J. Am. Chem. Soc., 70, 1982 (1948).

Skraup had suggested originally that the aromatic amine condensed with acrolein to form a Schiff base (VI), but this cannot be correct. If it were, β -methylacrolein (crotonaldehyde) should yield as an intermediate the Schiff base VII, which on ring closure would give 4-methyl-

quinoline (lepidine). The product, however, is 2-methylquinoline (quinaldine), and therefore the intermediate must be the β -arylamino-aldehyde VIII or a Schiff base derived from it.⁹

SCOPE AND LIMITATIONS

The Skraup reaction is of great general utility and has been applied to many aromatic amines. The only amines that fail to give the desired quinolines are those having substituents too reactive to withstand the drastic conditions, e.g., labile substituents such as acetyl, cyano, methoxyl, and fluoro. p-Aminoacetophenone, 2-cyano-5-methylaniline, 2-methoxyaniline, 3-nitro-4,5-dimethoxyaniline, 2-nitro-4-methoxy-5-fluoroaniline, and 3-nitro-4-aminoveratrole 4 fail to give the corresponding quinoline derivatives because the substituents are either degraded or hydrolyzed by the hot, strong sulfuric acid used in the reaction. The hydrolytic action of the sulfuric acid can be minimized by reducing the reaction time from the usual several hours to a few minutes. With a reaction time of one and one-half minutes 8-nitro-5,6-dimethoxyquinoline was prepared from 2-nitro-4,5-dimethoxyaniline in 40% yield. 13

The original Skraup synthesis has been extended to include the preparation of quinolines substituted in the pyridine ring through the

⁹ Manske, Chem. Revs., 30, 113 (1942).

¹⁰ Berend and Thomas, Ber., 25, 2548 (1892).

¹¹ v. Jakubowski, Ber., 43, 3026 (1910).

¹² Kaslow and Raymond, J. Am. Chem. Soc., 68, 1102 (1946).

¹³ Elderfield, Gensler, Williamson, Griffing, Kupchan, Maynard, Kreysa, and Wright, J. Am. Chem. Soc., 68, 1584 (1946).

¹⁴ Frisch, Silverman, and Bogert, J. Am. Chem. Soc., 65, 2432 (1943).

use of α,β -unsaturated aldehydes and ketones. 2-Methylquinolines (X) are obtained in high yield by adding β -methylacrolein (crotonaldehyde) (IX),15 its diacetate,15 or 1,1,3-trimethoxybutane 16 to a stirred mixture

$$\begin{array}{c} CHO \\ + CH \\ NH_2 & CHCH_3 \\ IX & X \end{array} \rightarrow \begin{array}{c} CH_3 \\ N \\ X \end{array}$$

of sulfuric acid, an oxidant, and an aromatic amine at such a rate that violent reaction is avoided. 2-Arylquinolines are prepared similarly by employing β-phenylacrolein (cinnamaldehyde) in place of crotonaldehyde. 17,18,19 The use of an α-substituted acrolein (XI) 8,15,20 or a 2-substituted glycerol 21,22,23 as an addend in the Skraup reaction results in a quinoline substituted in the 3 position (XII, R = methyl, aryl, or halogen). The acetal, the diacetate, or the dipropionate of the α substituted acrolein is often preferred in order to avoid the polymeri-

$$\begin{array}{c} \text{CHO} \\ \downarrow \\ \text{NH}_2 \\ \text{CH}_2 \\ \text{XI} \end{array} \rightarrow \begin{array}{c} \text{N} \\ \text{N} \\ \text{XII} \end{array}$$

zation of part of the aldehyde during the reaction. 15,20

While engaged in a study of antimalarial compounds, Campbell and co-workers 16,24-27 synthesized some 4-methylquinolines (XIV, R = methyl) by condensing methyl vinyl ketone (XIII, R = methyl) with aromatic amines under conditions somewhat milder than those used by Skraup. In view of the fact that α,β -unsaturated ketones such as XIII polymerize to some extent under the conditions of the reaction, it has

- ¹⁵ Utermohlen, J. Org. Chem., 8, 544 (1943).
- 16 Campbell, Helbing, and Kerwin, J. Am. Chem. Soc., 68, 1840 (1946). 17 Murmann, Monatsh., 25, 621 (1904).
- 18 Grimaux, Compt. rend., 96, 584 (1883).
- ¹⁹ Elderfield, Gensler, Bembry, Williamson, and Weisl, J. Am. Chem. Soc., 68, 1589 (1946).
 - ²⁰ Manske, Marion, and Leger, Can. J. Research, 20B, 133 (1942).
 - ²¹ Darzens and Meyer, Compt. rend., 198, 1428 (1934).
 - 22 Warren, J. Chem. Soc., 1936, 1366.
 - ²³ Brown and Dougherty, J. Am. Chem. Soc., 69, 2232 (1947).
 - ²⁴ Campbell and Schaffner, J. Am. Chem. Soc., 67, 86 (1945).
 - ²⁵ Campbell, Sommers, Kerwin, and Campbell, J. Am. Chem. Soc., 68, 1851 (1946).
 - ²² Campbell, Sommers, Kerwin, and Campbell, J. Am. Chem. Soc., 58, 1556 (1946). Campbell, Elderfield, Gensler, Sommere, Kremer, Kupchan, Tinker, Dressner, Romanek, and Campbell, J. Am. Chem. Soc., 69, 1465 (1947).

been found expedient to employ compounds that will yield the α,β -unsaturated ketones under these conditions. Thus β -ketobutanol, 20, 28, 29

$$\begin{array}{c|c} R \\ \downarrow \\ CO \\ + \downarrow \\ CH \\ CH_2 \\ \downarrow \\ CH_2 \\ \times III \end{array} \rightarrow \begin{array}{c} R \\ \\ N \end{array}$$

methyl β -chloroethyl ketone, ^{30,31,32} 4-methoxy-2-butanone, ²⁴ and 1,3,3-trimethoxybutane ^{24-27,33} when condensed with aniline all yield 4-methylquinoline, presumably via methyl vinyl ketone. 1-Aryl-3-chloropropan-1-ones are used for the preparation of 4-arylquinolines (XIV, R = phenyl). ^{30,34}

Aroquinolines

Amino derivatives of such fused systems as naphthalene, anthracene, phenanthrene, and pyrene undergo the Skraup reaction readily, and the resulting products are classed as aroquinolines. With 1-naphthylamine only one compound, benzo(h)quinoline (XV), is possible, ^{22, 35-41} but 2-naphthylamine might react with glycerol in two ways to produce a mixture of the two isomers, benzo(f)quinoline (XVI) and benzo(g)quinoline (XVII). The ring closure actually takes place in the 1 position of 2-naphthylamine, and benzo(f)quinoline (XVI) is the only product. ^{36, 42-48}

- ²⁸ Prill and Walter, Ger. pat. 505,320 [C. A., 26, 479 (1932)].
- ²⁹ I. G. Farbenindustrie A.G., Brit. pat. 308,365 [C. A., 24, 128 (1930)].
- 30 Kenner and Statham, Ber., 69, 16 (1936).
- ³¹ Schering-Kahlbaum A.G., Brit. pat. 283,577 [C. A., 22, 4132 (1928)].
- 32 Zöllner, U. S. pat. 1,804,045 [C. A., 25, 3668 (1931)].
- 33 Campbell and Kerwin, J. Am. Chem. Soc., 68, 1837 (1946).
- 34 Kenner and Statham, J. Chem. Soc., 1935, 299.
- 35 Skraup, Ber., 14, 1002 (1881).
- 36 Skraup, Ber., 15, 893 (1882); Monatsh., 3, 531 (1882).
- ³⁷ Farbwerke vorm Meister, Lucius, and Brüning, Ger. pat. 26,430 (1883) [Frdl., 1, 183 (1877–1887)].
 - 38 I. G. Farbenindustrie A.G., Fr. pat. 727,528 [C. A., 26, 5104 (1932)].
 - 39 Claus and Imhoff, J. prakt. Chem., [2] 57, 68 (1898).
 - ⁴⁰ Bamberger and Stettenheimer, Ber., 24, 2472 (1891).
 - 41 Schenkel and Schenkel, Helv. Chim. Acta, 27, 1456 (1944).
 - 42 Mikbailov, Novosti Tekhniki, 1940, No. 3-4, 51 [C. A., 34, 5847 (1940)].
 - ⁴³ Knueppel, Ber., 29, 703 (1896).
 - 44 Claus and Besseler, J. prakt. Chem., [2] 57, 49 (1898).
 - 45 Bamberger and Müller, Ber., 24, 2641 (1891).
 - 46 Clem and Hamilton, J. Am. Chem. Soc., 62, 2349 (1940).
- ⁴⁷ Sergeev, Byull. Lako-Krasochnol Prom., 1938, No. 2-3, 68; Khim. Referat. Zhur., 2, No. 1, 102 [C. A., 34, 1665 (1940)].
 - 48 Skraup and Cobenzl, Monatsh., 4, 436 (1883).

So strong is the tendency for ring closure to occur in the 1 position that a substituent such as halogen or nitro (but not methyl) in that position in 2-naphthylamine is eliminated. Thus 1-nitro-,49,50 1-bromo-,49,50 and 1-chloro-2-naphthylamine 51,52 when subjected to the Skraup reaction yield benzo(f)quinoline (XVI) alone or in admixture with the corresponding 10-substituted benzo(g)quinoline. In contrast to this, 5,6,7,8tetrahydro-2-naphthylamine undergoes the Skraup reaction to yield a mixture of 7,8,9,10-tetrahydrobenzo(f)quinoline and 6,7,8,9-tetrahydrobenzo(g)quinoline, with the latter predominating.53 amines that undergo this reaction are 1-, 2-, 3-, 4-, and 9-aminophenanthrene, 54, 55, 56 3-aminopyrene, 57 3-aminoacenaphthene, 58 1- and 2-aminoanthraquinone, 43, 59-64 and 2-aminofluorene. 55 Heterocyclic amines such as 3-aminopyridine, 66 2-aminothiophene, 67 and the aminobenzopyrones 68,69 do not withstand the drastic conditions well, and therefore the yields of the resulting quinoline derivatives in general are poor.

Aromatic diamines react with two moles of glycerol to give products known as phenanthrolines. The preparation of 1,7- (XVIII) 36,43,70,71,72

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49 Lellmann and Schmidt, Ber., 20, 3154 (1887).
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50 Huisgen, Ann., 559, 101 (1948).

⁵¹ Gerhardt and Hamilton, J. Am. Chem. Soc., 66, 479 (1944).

52 Clemo and Driver, J. Chem. Soc., 1945, 829. 53 v. Braun and Gruber, Ber., 55, 1710 (1922).

54 Herschmann, Ber., 41, 1998 (1908).

55 Cook and Thomson, J. Chem. Soc., 1945, 395.

55 Mosettig and Krueger, J. Org. Chem., 3, 317 (1938).

57 Vollmann, Becker, Corell, Streeck, and Langbein, Ann., 531, 1 (1937).

58 Zinke and Raith, Monatsh., 40, 271 (1919).

59 Delaby and Hiron, Bull. soc. chim. France, [4] 47, 227, 1395 (1930).

60 Majert, Ger. pat. 26,197 [Frdl., 1, 171 (1877-1887)].

61 Farbwerke vorm Meister, Lucius, and Brüning, Ger. pat. 189,234 [Frdl., 8, 1362 (1905-1907)].

⁶² Badische Anilin- und Sodafabrik, Ger. pat. 171,939 [Frdl., 8, 369 (1905–1907)].

⁶³ Schaarschmidt and Stahlschmidt, Ber., 45, 3452 (1912).

64 Graebe, Ann., 201, 333 (1880).

45 Hughes, Lions, and Wright, J. Proc. Roy. Soc. N. S. Wales, 71, 449 (1938) [C. A., 33, 609 (1939)].

66 Allen, Chem. Revs., 47, 275 (1950).

67 Steinkopf and Lützkendorf, Ann., 403, 45 (1914).

es Dey and Goswami, J. Chem. Soc., 115, 531 (1919).

es Dhar, J. Chem. Soc., 117, 1053 (1920).

⁷⁰ Druce, Chem. News, 119, 271 (1919) [C. A., 14, 535 (1920)].

71 Smith, J. Am. Chem. Soc., 52, 397 (1930).

7 Skraup and Vortmann, Monatsh., 3, 570 (1882); 4, 569 (1883).

and 4,7-phenanthroline (XIX) $^{36,70-74}$ from m- and p-phenylenediamine, respectively, offers no difficulties. Although some workers have reported failure of attempts to prepare 1,10-phenanthroline (XX) from o-phenylenediamine, 71,75 others have reported yields of 30-45%. 76,77 A

$$\begin{array}{c|c}
N_1 & 2 & 3 \\
\hline
0 & 10 & 4 \\
\hline
0 & 10 & 4$$

far better method for preparing 1,10-phenanthroline is to subject 8-aminoquinoline ^{71,78} or its derivatives ^{79,80} to the Skraup reaction. 8-Aminoquinolines are readily obtained from the corresponding onitroanilines by way of the 8-nitroquinolines. It is to be noted that 5- and 6-substituted 8-aminoquinolines yield identical phenanthroline derivatives. 4-Aminoquinolines ^{81,82} and 5-aminoisoquinolines ⁷⁵ undergo the Skraup reaction, but the yields are poor.

A double Skraup reaction also occurs with a diaminobiphenyl. A

good example is the preparation of 6,6'-biquinolyl (XXI) from 4,4'-diaminobiphenyl (benzidine).^{83, 84, 85} Another method for the preparation of biquinolyls is the Skraup synthesis with an anilinoquinoline, e.g., 4-methyl-2,6'-biquinolyl (XXII) from 2-p-anilino-4-methylquinoline.⁸⁶

- 73 Haskelberg, J. Am. Chem. Soc., 69, 1538 (1947).
- ⁷⁴ Douglas, Jacomb, and Kermack, J. Chem. Soc., 1947, 1659.
- 75 Misani and Bogert, J. Org. Chem., 10, 347 (1945).
- ⁷⁶ Halcrow and Kermack, J. Chem. Soc., 1945, 155.
- 7 Breckenridge and Singer, Can. J. Research, 25B, 583 (1947).
- 78 Smith and Gctz, Chem. Revs., 16, 113 (1935).
- 79 Richter and Smith, J. Am. Chem. Soc., 66, 396 (1944).
- 80 Burger, Bass, and Frederickson, J. Org. Chem., 9, 373 (1944).
- 81 Marckwald, Ann., 279, 20 (1894).
- ⁸² Lions and Ritchie, J. Proc. Roy. Soc. N. S. Wales, 74, 443 (1941) [C. A., 35, 4771 (1941)].
 - 83 Roser, Ber., 17, 1817, 2767 (1884).
 - 84 Ostermayer and Henrichsen, Ber., 17, 2444 (1884).
 - 85 Fischer, Monatsh., 5, 417 (1884).
 - 86 Fischer, Ber., 19, 1036 (1886).

5-Aminohydrindene also follows this rule, yielding a mixture of 6,7-trimethylene- and 5,6-trimethylene-quinoline in the ratio of 9:1.94

Application of the Skraup synthesis to 2-naphthylamine, $^{36,42-48}$ 2-aminofluorene, 66 and 2-aminophenanthrene 56 yields the angular isomers only, benzo(f)quinoline (XVI), 11-indeno(f, f)quinoline (XXIII), and naphtho(f, f)quinoline (XXIV), respectively. On the

$$\begin{array}{c|c} H_2 & 1 & 2 & 3 \\ \hline & 11 & & 4N \\ 8 & 7 & & 6 & 5 \\ \hline & & & & & & \\ XX111 & & & & & & \\ XX1V & & & & & \\ \end{array}$$

other hand, 5,6,7,8-tetrahydro-2-naphthylamine ⁵³ gives a mixture in which the linear isomer, 6,7,8,9-tetrahydrobenzo(g)quinoline, predominates. Like 2-naphthylamine, 6-aminoquinoline undergoes the Skraup ring closure in the 5 position to yield the angular isomer, 4,7-phenanthroline (XIX), exclusively. 5-Nitro- and 5-bromo-6-aminoquinoline also lose the 5 substituent on cyclization to form 4,7-phenanthroline. 5-Methyl-6-aminoquinoline retains its 5 substituent, and the product is 10-methyl-1,6-anthrazoline (XXV). ⁵⁰ 7-Aminoquinoline undergoes the Skraup reaction to yield the angular isomer, 1,7-phenan-

throline (XVIII), only. With 3-aminopyridine and 3-amino-2-chloropyridine the cyclization takes place at the 2 position to form only the linear compound, XXVI (1,5-naphthyridine), the halogen being eliminated in the latter case. The cyclization at the 4 position is evidently difficult since 3-amino-2,6-dimethylpyridine will not undergo the Skraup reaction. ⁶⁶ 3-Aminodibenzofuran produces a mixture of the two isomeric quinolines. ^{95, 96, 97}

Determination of the Identities of 5- and 7-Substituted Quinolines

In determining the identity of the two isomeric quinolines formed from *meta*-substituted anilines in the Skraup reaction, the synthesis of

⁹⁴ Lindner, Sellner, Hofmann, and Hager, Monatsh., 72, 335, 354 (1939),

⁹⁵ Mosettig and Robinson, J. Am. Chem. Soc., 57, 902 (1935).

⁹⁶ Kirkpatrick and Parker, J. Am. Chem. Soc., 57, 1123 (1935).

⁹⁷ Adams, Clark, Kornblum, and Wolff, J. Am. Chem. Soc., 66, 22 (1944).

one or both isomers by unambiguous methods is necessary. The most common method is to block one of the ortho positions of the metasubstituted aniline, subject it to the Skraup reaction, and then remove the blocking group from the resulting quinoline. To obtain 5-methylquinoline, 2-nitro-5-methylaniline 20 and 2-carboxy-5-methylaniline 11 were converted by means of the Skraup synthesis to 5-methyl-8-nitroand 5-methyl-8-carboxy-quinoline, respectively, and the 8 substituents then removed. In the same way toluene-2,3-diamine (2,3-diaminotoluene) 87 was converted to 7-methylquinoline by the Skraup synthesis followed by deamination of the resulting 7-methyl-8-aminoquinoline. Another method is to introduce further substituents into the two isomeric quinolines and then compare the products with compounds synthesized in an unequivocal way. Thus, the isomeric chloroquinolines obtained from m-chloroaniline were nitrated and the resulting products, 5-chloro-8-nitro- and 7-chloro-8-nitro-quinoline, proved to be identical with those obtained from 2-nitro-5-chloroaniline and 7-hydroxy-8-nitroquinoline. 93, 99, 100

The less common method for determining the identities of the 5- and 7-substituted quinolines is the synthesis of these compounds by an unambiguous method. In the Pfitzinger, Friedländer, Camps, and v. Niementowski quinoline syntheses, the hetero ring is formed by linking the ends of a two-carbon chain to the amino group and the ortho substituent in an ortho-substituted aniline. The preparation of 5- and 7-substituted quinolines by these methods is therefore unequivocal. These syntheses have been frequently used to establish the identity of the 5- and 7-isomeric quinolines obtained from a metasubstituted aniline in the syntheses of Doebner-Miller, Conrad-Limpach-Knorr, and Combes. They may also be employed in the identification of the products of the Skraup reaction. The Pfitzinger synthesis provides 5- and 7-substituted 4-carboxyquinolines which on decarboxylation should yield the desired reference compounds.

EXPERIMENTAL CONDITIONS

Control of the Reaction

The conditions under which the earlier Skraup syntheses were carried out often resulted in reactions of uncontrollable violence. The gradual addition of one of the reagents (glycerol or sulfuric acid) does not

²⁸ Price and Guthrie, J. Am. Chem. Soc., 68, 1592 (1946).

³⁰ Lutz, Bailey, Martin, and Salsbury, J. Am. Chem. Soc., 68, 1324 (1946).
¹³⁰ Claus and Junghanns, J. prakt. Chem., [2] 48, 254 (1893).

moderate the reaction satisfactorily, and the yields are poor. The modification of Clarke and Davis,¹⁰¹ the addition of ferrons sulfate, does regulate the reaction, presumably because the ferrous sulfate functions as an oxygen carrier and therefore the reaction is extended over a longer period of time. Further improvement has been achieved by the addition of acetic ¹⁰² or boric acid.¹⁰³ Manske, Leger, and Gallagher ¹⁰⁴ observed that the use of acetanilide in place of aniline in conjunction with ferrons sulfate and boroglyceric acid resulted in further moderation so that mole runs in 3- to 5-1. flasks could be carried out with perfect safety and increased yield. A British patent claims that the use of dilute sulfuric acid in the Skraup reaction eliminates violence and reduces the formation of tars.¹⁰⁵ Other workers ^{42,106,107,108} prefer strong sulfuric acid and avoid dilution during the reaction by removal of the water formed as an azeotrope with nitrobenzene.

Though the above modifications of the original Skraup synthesis have reduced the hazards of the reaction considerably, the violence was not reduced sufficiently to permit the preparation of quinolines on a commercial scale. It was discovered recently 109 that the mode of addition of the reactants is the most important factor in controlling the vigor of the reaction. When the mixture of the aromatic amine, suffuric acid, and glycerol kept at 80° is added in small portions to the reaction vessel containing the oxidizing agent, the reaction can be maintained easily at the required temperature and good yields can be obtained in large-scale production.

arsenic pentoxide,43 ferric oxide or sulfate,110 ferric chloride,24 stannic chloride, 70,111 chloropicrin, 112,113 o-nitrophenol, 20 and iodine. 114

EXPERIMENTAL PROCEDURES

The preparation of quinoline 101 in quantities of 255–275 g. with yields of 84-91%, and the preparation of 6-methoxy-8-nitroquinoline 115 in quantities of 460-540 g. with yields of 65-75%, are described in Organic Syntheses.

Quinoline.104 To 20 g. of powdered crystalline ferrous sulfate in a 5-1. flask there are added with shaking, in the order named, 77.6 g. of acetanilide, 42 g. of nitrobenzene, a solution of 35.5 g. of boric acid in 216 g. of glycerol, and 182 g. of concentrated sulfuric acid. The solution is then heated gently under a reflux condenser until it begins to simmer. Careful heating is continued for one-half hour, after which time the heat is increased for a further three hours.

The solution is then cooled slightly, 300 ml. of water is added, and the mixture is steam-distilled to remove the excess nitrobenzene (about 10 g.). The residual solution is cooled, and a solution of 340 g. of sodium hydroxide in 1 l. of water is added. The alkaline mixture is steamdistilled to remove the quinoline. After the quinoline layer is separated from the distillate, the aqueous layer is distilled to recover a small additional amount of quinoline.

To the combined quinoline layers is added 70 g. of concentrated sulfuric acid, and the resulting solution is diazotized at 8° with an excess of aqueous sodium nitrite (1-2 g. is sufficient). The diazotized solution is heated on the steam bath for thirty minutes, then steam-distilled to remove volatile impurities. A solution of 100 g. of sodium hydroxide in 400 ml. of water is added to the residual solution, and the mixture is again steam-distilled. The aqueous layer in the distillate is again concentrated as described above, and the quinoline is extracted from the combined distillates by means of benzene. Removal of the benzene, followed by distillation of the residue at 110-114°/14 mm. furnishes 67 g. (90%) of water-white quinoline.

3-Ethylquinoline. Into 165 g. of 20% oleum at 20-30°, 37 g. (0.3 mole) of nitrobenzene is run slowly and the mixture is heated with stirring to 60-70° over a period of approximately three hours. The

¹¹³ Barnett, Chem. News, 121, 205 (1920) [C. A., 15, 831 (1921)].

¹¹¹ Druce, Chem. News, 117, 346 (1918) [C. A., 13, 289 (1919)].

¹²² Gardner and Williams, Brit. pat. 198,462 [C. A., 17, 3880 (1923)].

¹¹³ Kaulmann and Hüssy, Ber., 41, 1735 (1908).

¹¹⁴ Hewitt and Trustham, U. S. pat. 2,358,162 [C. A., 39, 1421 (1945)]. 23 Mosher, Yanko, and Whitmore, Org. Syntheses, 27, 48 (1947).

mixture is maintained at this temperature for an additional six to eight hours until a sample is completely soluble in water. This mixture of nitrobenzenesulfonic acid and sulfuric acid, which is termed the "sulfo mix," is poured into 50 ml. of water in a 1-l. three-necked flask, equipped with a short still head and variable-length finger condenser, a dropping funnel, a thermometer, and a stainless steel sweep stirrer. This dilutes the sulfuric acid to a concentration of 75%. With stirring, 47 g. of aniline (0.5 mole) is added; the aniline sulfate soon dissolves in the acid mixture.

The whole is heated to 125° in an oil bath, and 93 g. (0.5 mole) of α-ethylacrolein diacetate is added dropwise with stirring; the addition is momentarily stopped if the reaction becomes violent. Both during and after the addition of the acrolein acetate, the mixture is heated and stirred (stirring is momentarily stopped if excessive foaming occurs); meanwhile, the finger condenser is gradually moved up, so that a slow, steady distillation of water and acetic acid takes place. In about three hours the oil-bath temperature has been allowed to rise to 175°, about 50 ml. of distillate has come over, and distillation has almost ceased. The reaction mixture is partially cooled, poured onto about 500 g. of ice, and neutralized with concentrated sodium hydroxide solution. The crude product is removed by steam distillation, preferably with superheated steam. The 3-ethylquinoline is separated from the distillate, with the aid of carbon tetrachloride extraction. Fractionation of the solvent-quinoline mixture gives 42.5 g. (54%) of pure 3-ethylquinoline, b.p. $265-266^{\circ}$; $n_D^{20} = 1.5988$.

4-Methyl-6-methoxy-8-nitroquinoline.27 A mixture of 170 g. of arsenic acid, 50 ml. of water, 168 g. (1.0 mole) of m-nitro-p-anisidine, and 280 g. of concentrated sulfuric acid is placed in a 1-l. flask fitted with stirrer, dropping funnel, and condenser set for downward distillation. The mixture is heated in an oil bath at 110-115° while 148 g. (1.0 mole) of 1,3,3-trimethoxybutane is added dropwise in the course of two and a half hours. The mixture is stirred at 115-125° for an additional two hours while methanol distils. It is then poured into 1 l. of water, filtered, and the filtrate diluted successively to 3 and 6 l., filtering after each dilution. The precipitates (mostly tars) are discarded. The final filtrate is made basic with aqueous ammonia, and the reddish precipitate is collected and dried; the yield of crude product melting at 158-160° is about 168 g. This material is dissolved in 2-2.5 l. of 10% hydrochloric acid and the solution heated on the steam bath for fifteen minutes with Norit, then filtered. The cooled solution is neutralized with aqueous ammonia, and the dried precipitate recrystallized from 2-2.5 l. of ethyl acetate, using Norit. The mother liquors from the first crop are concentrated to 500 ml. to give a second crop. The total yield of material melting at 169-171° or higher is 130 g. (55-60%).

Separation of the Mixture of 3,7- and 3,5-Dimethylquinoline.²⁰ The mixture is prepared from *m*-toluidine and α-methylacrolein. After distillation of the mixture most of the pale greenish distillate crystallizes. The oil is drained off, and the solid 3,7-dimethylquinoline is crystallized twice from purified hexane; m.p. 80°. The oily mixture from which the solid base has crystallized is dissolved in hot dilute perchloric acid and cooled. The precipitate is collected, washed with cold water, and recrystallized from boiling water to obtain the pure perchlorate of 3,5-dimethylquinoline as brilliant colorless prisms, m.p. 216°. The 3,7-dimethylquinoline regenerated from the filtrate crystallizes at once and, after being pressed on filter paper, melts at 78°.

TABULAR SURVEY OF QUINOLINES PREPARED BY THE SKRAUP SYNTHESIS

In the tables that follow are listed the quinolines prepared by the Skraup reaction through August, 1951. Within each table the quinolines are listed according to the substituents present in the following sequence: halogen; nitro; hydroxy, alkoxy, aryloxy, and RCO₂—; sulfurcontaining groups; amino; cyano; carbonyl; carboxyl; alkyl; aryl; heterocyclic. A substance containing more than one of the above groups is listed according to the group lowest in the list. Thus a 5-nitro-8-methyl-quinoline would follow 5,8-dicarboxyquinoline and would precede 5,8-dimethylquinoline.

TABLE I

QUINOLINES

	S.	*11	101,	106,	114,	ŊΙ	14 T I	11.0			96I			VO					that	orran
	References		35, 42, 43, 70,	102, 103, 106, 107, 108, 110	111, 112,	4, 5	104 135	0	184	87, 98, 187, 1	38, 145, 177,	87, 98, 187, 196	195, 196, 291	87, 190, 193	7, 177	87, 190, 193	190, 197, 213	87, 228, 229	is taken from	***
Viold	%		84-91			20	- 80 -	•	86	22	62	89	1	35	89	35	1	59	ld reported	•
Reactants	Second Component	A. Quinoline and Monosubstituted Quinolines	Glycerol			Acrolein	Glycerol	i	Glycerol	Glycerol	Glycerol	Glycerol.	Glysonsi	Glycerol Glycerol	Classical	Clycetol	Classes	City cel 01	116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that	
	ine Aniline	A. Quinoline	Aniline			Λ niline $N_{\perp}\Lambda$ oot N_{\parallel}	N-Allyl-	Ē	p-Huoro-	m-Chloro		o-Chlore-	m-Bromo-	p-Bromo-	m-Bromo-	o-Bromo-	m-Nitro		m	-
	Quinoline		Quinoline *			30.00		A Fluore	5-Chloro-	6-Chloro	7-Chloro-	8-Chloro-	5-Bromo-	6-Bromo-	7-Bromo-	8-Bromo-	5-Nitro-	Note: Defend	reference.	* Oningline by

reference.

* Quinoline has been obtained in 5% yield from phenylhydroxylamine and glycerol,148 and in traces from azoxybenzene and

TABLE I-Continued

	References	7. C. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	7, 45, 119, 191, 228, 238, 284	43, 87, 228, 229, 234, 235, 278			36, 38, 87, 234	226, 227	7, 12, 18, 90, 165,	150, 167 27 169 170	104, 172, 173, 174	88, 168	89	87, 250	37, 43, 299	241			
	$_{\%}^{\rm Yield}$		20	14	55	<u> </u>	46	2	99	7	27	53	1 3	21	1	30	3 6	77	
TVBIE I	Quinolines Reactants	Aniline Second Component	A. Quinoline and Monosubshinted Lumoring Glycerol		m-Nitro-			m-Hydroxy- Glycerol	o-Hydroxy-	p-Methoxy-		m-intentoxy- Glycerol		p-Phenoxy-		— p-sulfonic acid (sulfaminc Gryceron	—p-sulfonamide (sulfanil- Glycerol	p-Acctaminophenyl methyl Glycerol	. sulfone
		Quinoline		6-Nitro-	7-Nitro-	P.Nitro	5-Hydroxy-	6-Hydroxy-	8-Hydroxy-	A D Call come	0-Intentory	7-Methoxy-	8-Methoxy-	6-Phenoxy-	5-sulfonic acid	-6-sulfonie acid	-6-sulfonamide	s mothyl cultone	(6-SO ₂ CH ₃)

_		# cut + cut	HOIL WIE
59 30 21 17, 18 22 30 104, 177	201 30 30	178 30 220 220 220 220 220 220 220 220 220	TOWER CI
Poor 40 40 31 12 53 53	45 45 37	Hoor 60	**************************************
HOCH(C,H ₃)CHOHCH ₂ OH CICH ₂ CH ₂ CCC,H ₃ C ₂ H ₅ OCH ₂ C(C,H ₃ -is ₉)(OH)CH ₂ OC ₂ H ₅ C ₄ H ₅ CH=CHCHO C ₂ H ₅ OCH ₂ C(OH)(C ₆ H ₅)CH ₂ OC ₂ H ₅ CICH ₂ CH ₂ COC ₆ H ₅ Glycerol	Glyceld Glycerd ClCH ₂ CH ₂ COC ₆ H ₄ CH ₃ -p CH ₃ O CH ₃ O	Glycerol — 178 ClCH₂CH₂COC₁0H₁(β) 41 30 Glycerol — 220 Glycerol — 219 Glycerol — 219 Glycerol — 219 Glycerol — 219 Glycerol — 220 Glycerol — 219 Glycerol — 229 Glycerol — 219 Glycerol — 219 Glycerol — 220	
Aniline Aniline Aniline Aniline Aniline Aniline P-Phenyl-	p-t neny-rav-acetyr- o-Phenyl- Aniline Aniline	2,7, 2	
2-Butyl- 4-Butyl- 3-Isobutyl- 2-Phenyl- 3-Phenyl- 4-Phenyl-	8-Phenyl- 4-p-Tolyl- 4-(3-Methyl-6-methoxy- phenyl)-	6-Diphenylmethyl- 4-(6-Naphthyl)- 4-(6-Naphthyl)- 6-c-Pyridyl- 6-2-Pyridyl- 6-3-Pyridyl- 6-7-Pyridyl- 6-7-Pyridyl- 7-c-Pyridyl- 8-c-Pyridyl- 8-2-Pyridyl- 8-2-Pyridyl- 8-3-Pyridyl- 8-7-Pyridyl- 8-7-Pyridyl- 6-2,6-Dimethyl-4-pyridyl)- 6-(2,6-Dimethyl-4-pyridyl)- 6-(2,6-Dimethyl-4-pyridyl)- 6-(2,6-Dimethyl-4-pyridyl)- 6-(2,6-Dimethyl-4-pyridyl)- 6-(2,6-Dimethyl-4-pyridyl)- 6-Piperidylmethyl- 7-(2-Benzimidazolyl)- 8-Piperidylmethyl- 8-Piperidylmethyl- 8-Piperidylmethyl- 8-Piperidylmethyl- 9-Piperidylmethyl-	iererice.

TABLE I-Continue	QUINOLINES
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				ORG	ANIC	RE.	ACTI	ONS	5					
	References	30, 32	20, 87, 153, 154, 155	20, 35, 42, 70, 112, 151, 152	20, 35, 42, 43, 111,	112, 155 156 263	264 166, 263	264	290 59	21, 22, 23	15 $30, 31, 32$	20 158	59 23	30 163
	Yield %	40	Poor	46	67	; «	۱ ۾	3 3	39 Poor	2 4	54 40	92 50	Poor 15	40 35
3	Renctants Second Component	A. Quinoline and Monosubstituted Quinolines—Continued	ClOH2COLES Glycerol	Glycerol	Glycerol	Glycerol	Glycerol Glycerol	Glyeerol Glyeerol	Gyerol Gyerol HOCH(C,H.)CHOHCH2OH	CH-COCH-COCH5)OHCH2OC2H5 CH-COCH-COCH0	CH2=C(C2H)(CCOCH3)2 CH2=C(C2H)(CCOCH3)2 CH2 CH2COC3H	Glyeerol Glyeerol	HOCH(C3H,)CHOHCH,OH	CICH, COC, H, CICH, CH, COC, H, Glycerol
		Anilino A. Quinoline and	Aniline m-Methyl-	p-Methyl-	m-Methyl-	o-Methyl-	#-F3C-	25 C	p-Carboxymethyl-	Aniline Aniline	Aniline Aniline	Aniline m -Ethyl-	o-Ethyl- Aniline	Aniline Aniline o-Propyl-
		Quinoline	4-Methyl-(Cont'd.)	6-Methyl-	7-Methyl-	8-Methyl-	5-Trifluoromethyl-	6-Triffuoromethyl-7-Triffuoromethyl-	8-Trifluoromethyl- 6-Carboxymethyl-	2-Ethyl- 3-Ethyl-		4-Ethyl- 7-Ethyl-	8-Ethyl- 2-Propyl-	3-Propyl- 4-Propyl- 8-Propyl-

294 is taken from that	58 sized, the yield reported	Where one reference is italicized, the yield reported is taken from that	94-98.
122 249 <i>91</i> , 93, 307 294	28 30 83 83 88	Glycerol Glycerol Glycerol Glycerol	
321 115, <i>117</i> , 118, 239 123	68 71 70	CH2=C(Br)CHU Glycerol Glycerol	2-Nitro-4-ethoxy- poxy)- 2-Nitro-4-(\gamma-phthalimido- nronoxy)-
7	09	Acrolein	2-Nitro-4-methoxy-
117, 118, 119, 120, 232, 248,			
	192	Glycerol	2-Nitro-4-methoxy-
7	29	Acrolein	4-Nitro-2-methoxy-
120, 263, 266, 284	28	Glycerol	4-Nitro-2-methoxy-
7, 173	59	Acrolein	5-Nitro-2-methoxy-
297	-	Glycerol	3-Nitro-4-methoxy-
92	1	Glycerol	3-Bromo-4-ethoxy-
92	1	Glycerol	2-Bromo-4-methoxy-
92	58	Glycerol	3-Bromo-4-methoxy-
270	40	Glycerol	5-Chloro-2-methoxy-
134	1	Glycerol	2,4-Dihydroxy-
80, 255	2	Glycerol	5-Nitro-2-hydroxy-
286	50	Glycerol	4-Chloro-2-hydroxy-
218, 231, 270	35	Glycerol	5-Chloro-2-hydroxy-
218	. 30	Glycerol	5-Chloro-4-hydroxy-
113	1	Glycerol	2,3-Dinitro-
113, 177, 308	63	Glycerol	2,4-Dinitro-
113	1	Glycerol	3,5-Dinitro-

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	Yield % References		$\begin{array}{ccc} 50 & 221, 224 \\ 15 & 223 \end{array}$	00-00	_	35 38, <i>269</i>	130 130		130		46 184 20 99 196	50 99, 196		90 79, 115, 120, 247	55 93, 150 196	38.	•	60 151 40 113	
	χ,	•		ō	6		<u>'</u>	5											
QUINOLINES	Renetants	Second Component Anilino Anilino and Monosubstituted Quinolines—Continued	p-(2-Benzimidazolyl)- Glycerol c-(2-Benzimidazolyl)- Glycerol p-(6-Methyl-2-benzo-thiazolyl)-	B. Disubstituted Quinolines			2,4-Dichloro- 3.4 Dibromo-			3,4-Dibromo-Glycerol	2,4-Dibromo-Glyeerol	3-Chloro-4-nitro-		4-Chloro-3-ntro-Glycerol		2.Chloro-5-nitro-	2-Chloro-4-nitro-Glycerol		3,4-Dinitro-
		Quinolino	6-(2-Benzimidazolyl)- 8-(2-Benzimidazolyl)- (1-(6-Methyl-2-benzo-	thiazoiyi)-	t r	5,7-Dichloro	6,8-Dichloro-	5,6-Dibromo-	5,7-Dibromo-	6,7-Dibromo-	6,8-Dibronto-	6-Fluoro-8-nitro-	5-Chloro-8-nitro-	6-Chloro-7-nitro-	6-Chloro-8-nitro-	8-Chloro-5-nitro-	8-Chloro-6-nitro-	6-Bromo-8-nitro- 8-Bromo-6-nitro-	5,6-Dinitro-

THE SKRAUP SYNTHESIS O	F QUINOLINES	79
23 23 44	120, 232, 248, 284 7 321 115, 117, 118, 239 123 249 91, 93, 307 294	is taken from that
63 30 30 30 50 40 40 40 50 50 67 67	60 68 71 70 28 30 83 83	ized, the yield reported
Glycerol	Acrolein CH ₂ =C(Br)CHO Glycerol Glycerol Glycerol Glycerol Glycerol Glycerol	where one reference is italicized, the yield reported is taken from that
3,5-Dinitro- 2,4-Dinitro- 2,3-Dinitro- 5-Chloro-4-hydroxy- 5-Chloro-2-hydroxy- 5-Nitro-2-hydroxy- 2,4-Dihydroxy- 5-Chloro-2-methoxy- 3-Bromo-4-methoxy- 3-Bromo-4-ethoxy- 3-Bromo-4-ethoxy- 3-Nitro-4-methoxy- 4-Nitro-2-methoxy- 5-Nitro-2-methoxy- 3-Nitro-4-methoxy- 5-Nitro-2-methoxy- 5-Nitro-2-methoxy- 5-Nitro-2-methoxy- 5-Nitro-2-methoxy- 5-Nitro-2-methoxy- 5-Nitro-2-methoxy- 5-Nitro-4-methoxy- 5-Nitro-4-methoxy- 5-Nitro-4-methoxy- 5-Nitro-4-methoxy-	2-Nitro-4-methoxy- 2-Nitro-4-ethoxy- 2-Nitro-4-(γ -phthalimi propoxy)- 2-Nitro-4-butoxy- 2-Nitro-4-phenoxy- 3,4-Dimethoxy- 2,3-Dimethoxy-	
5,7-Dinitro- 6,8-Dinitro- 7,8-Dinitro- 5-Chloro-8-hydroxy- 6-Chloro-8-hydroxy- 6,8-Dihydroxy- 5-Nitro-8-methoxy- 7-Bromo-6-methoxy- 7-Bromo-6-methoxy- 5-Nitro-6-methoxy- 5-Nitro-8-methoxy- 5-Nitro-8-methoxy- 6-Nitro-8-methoxy- 8-Nitro-6-methoxy- 8-Nitro-6-methoxy- 8-Nitro-6-methoxy- 8-Nitro-6-methoxy-	ethoxy- (Y-aminopi butoxy- phenoxy- hoxy- hoxy-	reference.

I-Continued
ABLE

	References	88, 307	88, 89 89	232	258	87	126	87 256, 296	153	25 15, 244	32, <i>33</i> 161	15 16	15 25	87 151, 244
,	Yield %	Poor	62	1 1	88	83	65	50	1	55	55	90	52 67	74
Quinolines	Reactants Second Component	niline B. Disubstituted Quinolines—Continued	Glycerol Glycerol	Glycerol	Glycerol	Glycerol		y- Glycerol Glycerol			CH3CH=CHCH(OCOCH3)2 CH3C(OCH3)2CH2CH2OCH3	Glycerol Crotonaldehyde GH CH(OCH,)CH,CH(OCH3)2	CH2=C(CH3)CH(OCOCH3)2 CH_C(OCH3)CH(OCOCH3	Glycerol CH ₂ CH=CHCHO
		Anılıne B. Disubstit	3,4-Methylenedioxy- 3,4-Ethylenedioxy-	3,4-Phenylenedioxy-4-Methoxy-2-(1-diethyl-	amino-1-pentylammo/- 2-Chloro-5-sulfonic acid	2-Chloro-5-carboxy- 2-Bromo-5-carboxy-	5-Nitro-2-carboxy-	2-Hydroxy-4-carbomethoxy-	5-Methoxy-2-carboxy-4Methoxy-2-carbomethoxy-	2,5-Dicarboxy-	p-Chloro-	p-Cilion 2-Methyl-4-chloro- m-Chloro-	m-Chloro- m -Chloro-	m-Chloro- 3-Chloro-2-methyl- o-Chloro-
		Quinolino	6,7-Methylenedioxy-	6,7-Ethylenedloxy- 6,7-Phenylenedloxy- e x cohovy-8-(1-diethyl-	nnino-f-pentylmino)-	S-Chloro-5-earboxy-	5-Nitro-8-earboxy-	8-11ydroxy-5-curboxy-	5-Methoxy-8-carboxy-	6-Methoxy-8-carboxy- 5,8-Dicarboxy-	5-Chloro-4-methyl- 6-Chloro-2-methyl-	6-Chloro-4-methyt- 6-Chloro-8-methyt- 7 Chloro-2-methyt-	7.Chlore-3-methyl-	7-Chloro-4-methyl- 7-Chloro-8-methyl- 8-Chloro-2-methyl-

THE SKRAUP SYNTHESIS OF QUINOLINES	81
24 121, 273 275 20 161, 277 275 309 280, 281 15 15 15 15 16 280 287 319 319 319 319 319 319 319 319	d is taken from that
82 11 82 82 83 7 14 15 8 14 15 15 16 16 16 16 16 16	reporte
CH ₃ C(OCH ₃) ₂ CH ₂ CH ₂ OCH ₃ Glycerol	where one reference is italicized, the yield reported is taken from that
o-Chloro-2-methyl-5-Bromo-2-methyl-7-Bromo-2-methyl-3-Bromo-2-methyl-2-Bromo-2-methyl-2-Bromo-2-methyl-2-Bromo-4-methyl-5-Nitro-2-methyl-7-Nitro-2-methyl-4-Nitro-2-methyl-4-Nitro-2-methyl-3-Nitro-2-methyl-3-Nitro-2-methyl-2-Nitro-2-methyl-2-Nitro-3-methyl-3-Nitro-3-methyl-3-Nit	
8-Chloro-4-methyl- 8-Ciloro-5-methyl- 5-Bromo-8-methyl- 6-Bromo-8-methyl- 7-Bromo-6-methyl- 5-(or 7-)Nitro-6-methyl- 6-Nitro-8-methyl- 6-Nitro-3-methyl- 6-Nitro-3-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 8-Nitro-1-methyl- 8-Nitro-1-methyl- 8-Nitro-3-methyl- 8-Nitro-1-methyl- 8-Nithoxy-2-methyl- 8-Nitro-1-methyl- 8-Nithoxy-1-methyl- 6-Nithoxy-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 6-Nitro-1-methyl- 8-Nitro-1-methyl- 8-Nitro-1-methy	reference,

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75,214	γ_o References		83 292 83 293 83 287, 293 — 136	22 87	$\frac{231,32}{70}$		008 00 01	62 24 62 15	, , , ,		42 20 - 20 50 50	$\begin{array}{ccc} 12 & 20 \\ 54 & 15 \\ 65 & 15, 20 \end{array}$	
Quinolines	Renetants Second Component	n. Disubstituted Quinolines—Continued	Glyeerol Glyeerol Glyeerol Glyeerol	CH3CH=CHCHC	CH3CH=CHCHO CH3COCH2CH2CI	Glycerol Glycerol	•	CH,COCH,CHOHCH, CH,COCH=CHCH,	CHICH CHCH(OCOC2H5)2	CH ₃ CH=CHCH(OCOCH ₃) ₂ CH ₃ CH=CHCH(OCOCH ₃) ₂	CH3CHCCH3CH2OH CH3COCH(CH3)CH2OH	CH2=C(CH3)CHC CH2=C(CH3)CH(OCOCH3)2 CH2=C(CH3)CH(OCOCH3)2	CH2=C(CH3)CH(OCOCH3)2 CH2=C(CH3)CH(OCOCH3)2
	o ilino	R. Disubstitu	4-Methyl—3-sulfonic acid 4-Methyl—2-sulfonic acid 2-Methyl—5-sulfonic acid 2-Methyl—5-sulfonic acid	p-Arsonamino-(arsanilic acid)	2-Amino-3-methyl- p -Carboxy-	o-Carboxy- 2-Cyano-5-methyl- 6-Crond Amethyl-	4-sulfonie aeid (sulfanilie	acid) Aniline	$_{\tau\text{-Methyl-}}$	m-Methyl- m -Methyl-	o-Methyl- Aniline	m-Methyl- p -Methyl- m -Methyl- m -Methyl-	m-Methyl- o -Methyl-
		Quinoline	6-Methyl—7-sulfonic acid 6-Methyl—8-sulfonic acid 8-Methyl—5-sulfonic acid	8-Methyl—6-sulfonie acid 6-Arsonamino-2-methyl-	8-Amino-7-methyl- 6-Carboxy-2-methyl-	8-Carboxy-4-methyl- 8-Carboxy-5-methyl-	8-Carboxy-6-methyl-	2.4-Dimethyl-	4,1-1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,	2,6-Dimethyl- 2,7-Dimethyl-	2,8-Dimethyl-	3,5-Dimethyl- 3,5-Dimethyl- 3,6-Dimethyl-	3,7-Dimethyl- 3,8-Dimethyl-

THE SKRAUP SYNTHESIS OF QUINOLINES	8
18 20 65 24 — 29 Poor 20 39 20, 253 — 20, 112, 179 67 20, 245 70 20, 245 70 20, 245 70 20, 245 70 20, 245 70 20, 245 71 28 32 15 72 257, 276 — 268 15 19 40 282 40 282 40 282 50 282 40 282 61 125 61 125 61 125 61 125 61 125 61 282 62 282 63 282 64 282 65 61 66 282 67 282 68 124 69 69 — 90 — 90	reported is taken from that
CH2—C(CH3)CHO CH3COCH—CH2 CH3COCH2CH2 CH3COCH2CH2 CH3COCH2CH2 CH3COCH2CH2 CH3COCH2CH2 CH3cerol CH3cerol CH2—C(C2H4)CH(OCOCH3)2 CH2—C(C2H4)CHO CH3CH2COCH2CH2 CH3CH3CH2COCH2CH2 CH3CH3CH2COCH2CH2 CH3CH3CH2COCH2CH2 CH3CH3CH2COCH2CH2 CH3CH3CH2COCH2CH2 CH3CH3CH3CH2 CH3CH3CH3CH2 CH3CH3CH3CH3CH3 CH3CACH3CH2CH3CH3 CH3CACH3CH2CH3CH3 CH3CACH3CH3CH3CH3 CH3CACH3CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3C	The second of the second of the second reference is italicized, the yield reported is taken from that
o-Methyl- p-Methyl- 3,4-Dimethyl- 3,5-Dimethyl- 3,5-Dimethyl- 2,5-Dimethyl- 2,4-Dimethyl- 2,3-Dimethyl- 2,3-Dimethyl- 2,3-Dimethyl- 2,3-Dimethyl- 2,3-Dimethyl- 2-Methyl- 2-Methyl- 2-Methyl- 2-Methyl- 2-Methoxy- 2-Methyl-5-isopropyl- 4-Methoxy-2-isoamyl- o-Nitro- o-Nitro- 2-Nitro-5-phenyl-N-acetyl- 2-Nitro-4-phenyl-N-acetyl- 2-Nitro-4-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-phenyl- 2-Hydroxy-5-pridyl- 2-Hydroxy-5-c-pyridyl- 4-Methoxy-3-c-pyridyl- 4-Methoxy-2-c-pyridyl- 4-Methoxy-2-c-pyridyl- 4-Methoxy-2-p-pyridyl- 4-Methoxy-2-c-pyridyl- 4-Methoxy-2-p-pyridyl-	The makes on Pp. 32-30. Willer
thyl-Contii thyl-hyl-hyl-hyl-hyl-hyl-hyl-hyl-hyl-hyl-	

Yield References $ec{arphi}_0$	90	60 183 60 183 40 183	88 88 I	50 50 50 51 52 53 53 53 53 54 54 60 60 60 60 60 60 60 60 60 60	
Renetants Second Component	ted Quin	4-Methoxy-2- γ -pyridyl-Glyeerol 2-Methoxy-5- α -pyridyl-Glyeerol 5- ι -Dipyridyl-Glyeerol 2,5-Dipyridyl-Glyeerol 9.4-Dipyridyl-	C. Trisubstituted	3,4,5-Trichloro- 2,4,5-Trichloro- 2,4,5-Trichloro- 2,4,5-Trichloro- 3,5-Dichloro-2-hydroxy- 2-Nitro-4-bromo- 2-Nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-nitro-4-methoxy- 5-Chloro-2-methyl-5-nitro- 6lycerol	
	Quinoline	6-Methoxy-8-7-pyridyl-8-Methoxy-5-α-pyridyl-5-L-Butyl-8-pyridyl-†5-S-Dipyridyl-†	6,8-Dipyridyl- †	5,6,7-Trichloro- 5,6,8-Trichloro- 6,8-Dichloro-8-hydroxy- 3,7-Dichloro-8-hydroxy- 3,6-Dibromo-8-nitro- 5,6-Dibromo-8-nitro- 5,6-Dibromo-8-nitro-6-methoxy- 3-Ghloro-8-nitro-6-methoxy- 5-Fluoro-8-nitro-6-methoxy- 5-Chloro-8-nitro-6-methoxy- 5-Chloro-8-nitro-6-methoxy- 5-Dibromo-8-nitro-6-methoxy- 5-Dibromo-8-nitro-6-methoxy- 6-Chloro-8-nitro-6-methoxy- 6-Chloro-8-nitro-6-methyl- 6-Bromo-5-nitro-8-methyl- 8-Bromo-5-nitro-6-methyl- 6-Bromo-5-nitro-6-methyl- 6-Bromo-5-nitro-6-methyl- 6-Bromo-5-nitro-6-methyl-	

15 75 40 6, 13, 251 50 318 53 318 — 144, 307 — 267	- 267 5 252 0 26, 27 7 254		271 20, 317 317 20, 317 317 317 ed is taken from that
15 40 50 53 —	36 60 57	20	30 28 34 9 36 36
Glycerol Glycerol Acrolein Acrolein Glycerol	Glycerol CH ₃ CH=CHCHO CH ₃ OCH ₂ CH ₂ C(OCH ₃) ₂ CH ₃ Glycerol	Glycerol Glycerol Glycerol Glycerol Glycerol Glycerol	8-Nitro-3,5-dimethyl- 2-Nitro-4,6-dimethyl- CH ₂ COCH ₃)CH(OCOCH ₃)2 CH ₂ COCH ₃)CH(OCOCH ₃)2 CH ₂ COCH ₃ CH ₃ CH CH ₃ COCH C
5-Bromo-3,4-dimethoxy-2-Nitro-4,5-dimethoxy-2-Nitro-4,5-methylenedioxy-2-Nitro-4,5-cthylenedioxy-2,3,4-Trimethoxy-5-Bromo-4-methoxy-2-methyl-	3-Bromo-4-methoxy- 2-methyl- 2-Nitro-4-methoxy- 2-Nitro-4-methoxy- 5-mosthyl-	3-Amino-2-methyl—5-sulfonic Glycerol acid 3-Amino-5-carboxy- 2-methyl- 5-Nitro-2,4-dimethyl- 4-Nitro-2,5-dimethyl- 3-Nitro-2,3-dimethyl- 3-Nitro-2-methyl- Glycerol 3-Nitro-2-methyl- Glycerol 5-isopropyl-	Sylitro-3,5-dimethyl-2-Nitro-5-methyl-CH-2-Nitro-3,6-dimethyl-2-Nitro-4,methyl-CH-2-Nitro-4,fordimethyl-2-Nitro-4,fordimethyl-2-Nitro-4,fordimethyl-2-Nitro-4,fordimethyl-2-Nitro-4-methyl-CH-2-Nitro-4,fordimethyl-2-Nitro-4-methyl-CH-2-Nitro-4,fordimethyl-2-Nitro-4-methyl-CH-2-Nitro-4,fordimethyl-CH-2-Nitro-4-methyl-CH-2-Nitro
5-Bromo-6,7-dimethoxy-8-Nitro-5,6-dimethoxy-8-Nitro-5,6-methylenedioxy-8-Nitro-5,6-ethylenedioxy-6,7,8-Trimethoxy-5-Bromo-6-methoxy-8-methyl-	7-Bromo-6-methoxy- 8-methyl- 8-Nitro-6-methoxy-2-methyl- 8-Nitro-6-methoxy-4-methyl- 8-Nitro-6-methoxy-5-methyl-	7-Amino-8-methyl—5-sulfonic acid 7-Amino-5-carboxy- 8-methyl- 5-Nitro-6,8-dimethyl- 6-Nitro-5,8-dimethyl- 6-Nitro-7,8-dimethyl- 7-Nitro-5,9-dimethyl- 8-methyl- 8-methyl- 8-methyl- 8-methyl- 7-Nitro-7,9-dimethyl- 8-methyl-	8-Nitro-3,5-dimethyl- 8-Nitro-3,6-dimethyl- 8-Nitro-4,6-dimethyl- 8-Nitro-4,6-dimethyl- Note: References 116-322 are reference.

TABLE 1—Continued

			ORGAI	NIC I	REA	CTIO	S				
	References	20, <i>316</i> 122	140 141 141, 142	20 138, <i>139</i>	298 298	19 312	<i>2</i> 43	787	316	971	315
:	Xreld %	88	3111	27 Good	18 39	8 တွ	37	20	24	1	65
Quinolines	Reactants	Aniline C. Trisubstituted Quinolines—Continued		2,5-Dimethyl—3-sulfonie acid	2,4,5-Trimethyl-	4-Nitro-3-methyl- CH ₃ CH ₂	2.Nitro-4-methoxy- o Nitro-4-methoxy-	5-phenyl- 2-Nitro-4-phenyl- Columbia Columbia	D. Tetrasubstituted Quinolines	-dibromo-	N-acetyl- 2-Nitro-4-methyl- CH2=C(CH3)CH(OCOCH3)2 CH2=C(CH3)CH(OCOCH3)2
		Quinoline	8-Nitro-5, 6-dimethyl- 6-Methoxy-5, 7-dimethyl-	(FAmmo-3, Sumbourgers) 5,8-Dimethyl—G-sulfonic acid	3,4-7-Trinethyl-	5,0,5-1,1111cm; 6-Nitro-7-methyl-4-ethyl- 6-Nitro-8-methyl-4-ethyl-	g-Nitro-6-methoxy-2-phenyl-	S-Nitro-6-methoxy-5-phenyl-	8-Nitro-i, 6-dipheny i-	omought by a como-	8-Nitro-3,9,0-trimethyl-

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

TABLE II

Benzoquinolines

Benzo(h)quinoline

 N^{2}

A. Benzo(h)quinolines

\$\frac{10}{6}\$
Benzo(h)quinoline 7,8,9,10-Tetra- hydro- 6-Hydroxy-7,8,9,10- tetrahydro7-sulfonic acid -10-sulfonic acid 6-Methyl-

3-Ethyl-

6,7-Ace-

F	leactants		
1-Naphthylamine 1-Naphthylamine 5,6,7,8-Tetra- hydro-	Second Component Glycerol Glycerol	Yield % 25 —	References 35, 36, 39, 40 149
4-Hydroxy-5,6,7,8- tetrahydro-	Glycerol	25–30	260
—5-sulfonic acid —8-sulfonic acid 4-Methyl-	Glycerol Glycerol Glycerol	60	37, 38 41
1-Naphthylamine	OH C ₂ H ₅ OCH ₂ CCH ₂ OC ₂ H ₅	12	230
4,5-Ace-	C ₂ H ₅ Glycerol	35	58

B. Benzo (f) quinolines

1 2 3 N 8 2 6 5
Benzo(f)quinoline

N N	F			
Benzo())quinoline	2-Naphthylamino 2-Naphthylamine	Second Component Glycerol	Yield % 81	References 36, 42, 43, 44,
7.8,9,10-Tetra- hydro-	5,6,7,8-Tetra- hydro-	Glycerol	19	45, 46, 47, 48, 49, 216 53
10-Chloro- 10-Bromo-	1.8-Dichloro- 1-Chloro-8-bromo-	Glycerol Glycerol	15 38	283
8-Nitro- 10-Nitro- —8-sulfonic acid	6-Nitro- 8-Nitro- —6-sulfonic acid	Glycerol Glycerol Glycerol	34 34	52 46 46
-10-sulfonic acid 5-Carboxy-	-8-sulfonic acid 3-Carboxy-	Glycerol Glycerol	-	38 159
6-Carboxy- 1-Methyl- 3-Methyl-	4-Carboxy- 2-Naphthylamine 2-Naphthylamine	CH ² CH=CHCHO CH ² CH=CHCHO CH ³ CH ⁴ CH ⁴ CH ³ CH ⁴	29 — 53	242 157 24, 215
10-Hydroxy—8-sul- fonic acid	8-Hydroxy-6-sul- fonic acid	Glycerol	-	244 28, 162

Note: References 116-322 are listed on pp. 94-98. Where one reference is italioized, the field reported is taken from that reference.

TABLE II-Continued

Benzoquinolines

C. Benzo(g)quinolines

10 1 2	Re	eactants	Yield			
5 5	2-Naphthylamine	Second Component	%	References		
6.7.8.9-Tetrahydro-	5.6.7.8-Tetrahydro-	Glycerol	36	53		
10-Chloro-	1-Chloro-	Glycerol	34	51, 52		
10-Methyl-	1-Methyi-	Glycerol	25	50		
6.10-Dichloro-	1.5-Dichloro-	Glycerol		283		
5,10-Dichloro-	1,4-Dichloro-	Glycerol		283		
10-Chloro-6-bromo-	1-Chloro-5-bromo-	Glycerol		52		
10-Chloro-6-nitro-	1-Chloro-5-nitro-	Glycerol		51		
10-Chloro-7-nitro-	1-Chloro-6-nitro-	Glyecrol	4	51, 52		
10-Chloro-9-nitro-	1-Chloro-8-nitro-	Glycerol	12	51		
5,10-Diphenyl-	1,4-Diphenyl-	Glycerol	40-50	233		
9-Hydroxy-7-sul- fonic acid	8-Hydroxy—6-sul- fonic acid	Glycerol	~	162		

Note: References 116-322 are listed on pp. 94-98 Where one reference is italicized, the yield reported is taken from that reference.

TABLE III

BIQUINOLYLS

Biquinolyls are numbered to show the carbon atoms through which the two quinoline nuclei are joined; e.g., 2,7'-biquinolyl is

		Reactants		
Biquinolyl	Amine	Second Component	Yie	- Melet-
2,5'-	2-m-Aminophenyl- quinoline	Glycerol	% 21	311003
2,7′-	2-m-Aminophenyl- quinoline	Glycerol	30	189, 225
4,6'-	4-p-Arninophenyl- quinoline	Glycerol	-	188
4,7′-	4-m-Aminophenyl-	Glyccrol	_	188
6,6'-	4,4'-Diaminobiphenyl	Glycorol	80	83, 84, 85,
6,8'-	2,4'-Diaminobiphenyl	Glycerol		198
8,8'-	2,2'-Diaminobiphenyl	Glycorol	50	199
6-Methoxy-2,5'-	2-m-Aminophenyl- 6-methoxyquinoline	Glycerol	65 21	200 191
6-Methoxy-2,7'-	2-m-Aminophenyl- 6-methoxyquinoline	Glycerol	32	191
2'-(p-Nitrophenyl)- 2,6'-	2-p-Aminophenyl- quinoline	p-NO2C6I14CII=CIICHO	6	301
4-Methyl-2,6'-	2-p-Aminophenyl- 4-methylquinoline	Glycorol		36
8,8'-Dihydroxy-5,5'-	3,3'-Diamino-4,4'-dihy- droxybiphenyl	Glycerol	•-	-
5,5'-Dicarboxy-8,8'-	2,2'-Diamino-4,4'-di- carboxy biphenyl	Glycorol	A.	06
2,2'-Dimethyl-6,6'-	4.4'-Diaminobiphenyl	Crotomatt	19 20)5
5,5'-Dimethyl-8,8'-	2,2'-Diamino-4,4'-di- methylbiphenyl	Crotonaldehyde Glycerol	- 24s	
	-4 •		⁽⁴⁾ 200	1

Note: References 116-322 are listed on pp. 91-98. Where one releases is itslicked, the yield

TABLE V

PHENANTHROLINES

A. 1,10-Phenanthrolines

Phenanthroline	A. 1,10-Pheno	inthrolines		
<u> </u>				
N_1^2 3				
N. J.				
9 10		Reactants		
8 7 5	Amine	910	Yield	Refer
1.10.70 0. 11		Second Component	%	ences
1,10-Phenanthroline	o-Phenylenediamine	Glycerol	45	76
(5)6-Chloro-1,10-	8-Aminoquinoline 6-Chloro-8-aminoquinoline	Glycerol	40	71, 7
3-Bromo-1.10-	8-Amino-3-bromoquinoline	Glycerol	56	79
(5)6-Bromo-1,10-	5-Bromo-8-aminoquinoline	Glycerol Glycerol	20	316
(5)0 210110-1,10-	6-Bromo-8-aminoquinoline	Glycerol	40 46	76
(5)6-Nitro-1,10-	5-Nitro-8-aminoquinoline	Glycerol	40	79 76
2-Methyl-1,10-	2-Methyl-8-aminoquinoline	Glycerol	_	183
3-Methyl-1,10-	8-Aminoquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	- 6	315
4-Methyl-1,10-	8-Amino-4-methylquinoline	Glycerol	15	315
5(6)-Methyl-1,10-	8-Amino-6-methylquinoline	Glycerol	66	79
4-Phenyl-1,10-	8-Aminoquinoline	C ₆ H ₅ COCH ₂ CH ₂ Cl	15	282
- ,	4-Phenyl-8-aminoquinoline	Glycerol	Poor	282
5(6)-Phenyl-1,10-	6-Phenyl-8-aminoquinoline	Glycerol	20	282
3,5-Dibromo-1,10-	8-Amino-6-bromoquinoline	CH2=C(Br)CH(OCOCH3)2	1.4	316
3,6-Dibromo-1,10-	8-Amino-3,6-dibromoquinoline	Glycerol	28	316
3,8-Dibromo-1,10-	8-Amino-3-bromoquinoline	$CH_2 = C(B_r)CH(OCOCH_3)_2$	5	316
5,6-Dibromo-1,10-	8-Amino-5,6-dibromoquinoline	Glycerol	14	316
7-Chloro-3-methyl-1,10-	8-Amino-4-chloroquinoline	$CH_2 = C(CH_3)CH(OCOCH_3)_2$	3	315
2-Hydroxy-4-methyl-	8-Amino-2-hydroxy-4-methylquino-	Glycerol	20-30	80
1,10-	line	.		
2,9-Dimethyl-1,10-	8-Aminoquinaldine	Crotonaldehyde diacetate	7	315
3,4-Dimethyl-1,10- 3,5-Dimethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	Glycerol	22	271
0,0-Dimethyl-1,10-	8-Amino-6-methylquinoline	$CH_2 = C(CH_3)$ - $CH(OCOCH_3)_2$	4	317
3,6-Dimethyl-1,10-	8-Amino-3,6-dimethylquinoline	Glycerol	3	317
3,7-Dimethyl-1,10-	8-Amino-4-methylquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	2	315
3.8-Dimethyl-1.10-	8-Amino-3-methylquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	9	315
4.5-Dimethyl-1,10-	8-Amino-4,5-dimethylquinoline	Glycerol	24	317
4,6-Dimethyl-1,10-	8-Amino-4,6-dimethylquinoline	Glycerol	11	317
4,7-Dimethyl-1,10-	8-Amino-4-methylquinoline	CH ₃ OCH ₂ CH ₂ C(OCH ₃) ₂ CH ₃	7	315
5,6-Dimethyl-1,10-	8-Amino-5,6-dimethylquinoline	Glycerol	9	315
4,6-Diphenyl-1,10-	4,6-Diphenyl-8-aminoquinoline	Glycerol	10	282
4,7-Diphenyl-1,10-	4-Phenyl-8-aminoquinoline	C ₆ H ₆ COCH ₂ CH ₂ Cl	40	282
3,4,6-Trimethyl-1,10-	3,4,6-Trimethyl-8-aminoquinoline	Glycerol	19	271
3,4,7-Trimethyl-1,10- 3,4,8-Trimethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	CH ₃ COCH=CH ₂	31	271 271
3,5,6-Trihromo-1,10-	3,4-Dimethyl-8-aminoquinoline 8-Amino-3,5,6-tribromoquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂ Glycerol	9 27	316
3,5,6-Trimethyl-1,10-	8-Amino-5,6-dimethylquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	9	315
3,5,7-Trimethyl-1,10-	8-Amino-4,6-dimethylquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	15	317
3,5,8-Trimethyl-1,10-	8-Amino-3,5-dimethylquinoline	CH2=C(CH3)CH(OCOCH3)2	9	317
3,6,7-Trimethyl-1,10-	8-Amino-4,5-dimethylquinoline	CH2=C(CH3)CH(OCOCH3)2	2	317
4,5,7-Trimethyl-1,10-	8-Amino-4,6-dimethylquinoline	CH3COCH=CH2	1	317
3,5,6,8-Tetrabromo-1,10-	8-Amino-3,5,6-trihromoquinoline	CH2=C(Br)CH(OCOCH3)2	4	316
3,4,6,7-Tetramethyl-1,10-	3,4,6-Trimethyl-8-aminoquinoline	CH ₃ COCH=CH ₂	5	271
3,4,0,8-l'etramethyl-1,10-	3,4,6-Trimethyl-S-aminoquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	9	271
3.5.6.8.Tetramethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	CH ₃ COCH(CH ₃)CH ₂ OH	20 22	271 315
5,5,5,5-1 ett amet nyl-1,10-	8-Amino-3,5,6-trimethylquinoline	CH ₂ =C(CH ₃)CH(OCOCH ₃) ₂	44	010

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

TABLE IV

Compounds Containing Two or Three Quinoline Nuclei Separated by One or Two Carbon Atoms

D . 1 . 1	Danatanta	Yield	Refer-
$\mathbf{Product}$	Reactants	%	ences
6,6'-Diquinolylmethane	4,4'-Diaminodiphenylmethane + glycerol	19	202
6,6'-Diquinolyl ketone	4,4'-Diaminodiphenyl ketone + glycerol		203
Tri-(6-quinolyl)methane	Pararosaniline + glycerol		203
sym-6,6'-Diquinolyl- ethane	sym-4,4'-Diaminodiphenylethane + glycerol		204
sym-2,6'-Diquinolyl- ethylene	1-(p-Aminophenyl)-2-(2-quino- lyl)ethylene + glycerol		192

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

TABLE V

PHENANTHROLINES

A. 1,10-Phenanthrolines

Ph	enan	$_{ m thr}$	ol	ine

N. A				
9 10	F	leactants	Yield	Refer-
8 7 6 5	Amine	Second Component	7 ieia %	ences
1,10-Phenanthroline	o-Phenylenedismine	Glycerol	45	76
	8-Aminoquinoline	Glycerol	40	71, 78
(5)6-Chloro-1,10-	6-Chloro-8-aminoquinoline	Glycerol	56	79
3-Bromo-1,10-	8-Amino-3-bromoquinoline	Glycerol	20	316
(5)6-Bromo-1,10-	5-Bromo-8-aminoquinoline	Glycerol	40	76
	6-Bromo-8-aminoquinoline	Glycerol	46	79
(5)6-Nitro-1,10-	5-Nitro-8-aminoquinoline	Glycerol		76
2-Methyl-1,10-	2-Methyl-8-aminoquinoline	Glycerol		183
3-Methyl-1,10-	8-Aminoquinoline	CH2=C(CH3)CH(OCOCH3)2	6	315
4-Methyl-1,10-	8-Amino-4-methylquinoline	Glycerol	15	315
5(6)-Methyl-1,10-	8-Amino-6-methylquinoline	Glycerol	66	79
4-Phenyl-1,10-	8-Aminoquinoline	C6H5COCH2CH2Cl	15	282
	4-Phenyl-8-aminoquinoline	Glycerol	Poor	282
5(6)-Phenyl-1,10-	6-Phenyl-8-aminoquinoline	Glycerol	20	282
3,5-Dihromo-1,10-	8-Amino-6-hromoquinoline	CH2=C(Br)CH(OCOCH3)2	1.4	316
3,6-Dihromo-1,10-	8-Amino-3,6-dihromoquinoline	Glycerol	28	316
3,8-Dihromo-1,10-	8-Amino-3-hromoquinoline	CH ₂ =C(Br)CH(OCOCH ₃) ₂	5	316
5,6-Dihromo-1,10-	8-Amino-5,6-dihromoquinoline	Glycerol	14	316
7-Chloro-3-methyl-1,10-	8-Amino-4-chloroquinoline	CH2=C(CH3)CH(OCOCH3)2	3	315
2-Hydroxy-4-methyl-	8-Amino-2-hydroxy-4-methylquino-	Glycerol	20-30	80
1,10-	line			
2,9-Dimethyl-1,10-	8-Aminoquinaldine	Crotonaldehyde diacetate	7	315
3,4-Dimethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	Glycerol	22	271

3,5-Dihromo-1,10-	8-Amino-6-hromoquinoline	CH2=C(Br)CH(OCOCH3)2	1.4	316
3,6-Dihromo-1,10-	8-Amino-3,6-dihromoquinoline	Glycerol	28	316
3,8-Dihromo-1,10-	8-Amino-3-hromoquinoline	CH ₂ =C(Br)CH(OCOCH ₃) ₂	5	316
5,6-Dihromo-1,10-	8-Amino-5,6-dibromoquinoline	Glycerol	14	316
7-Chloro-3-methyl-1,10-	8-Amino-4-chloroquinoline	CH2=C(CH3)CH(OCOCH3)2	3	315
2-Hydroxy-4-methyl-	8-Amino-2-hydroxy-4-methylquino-	Glycerol	20-30	80
1,10-	line	•		
2,9-Dimethyl-1,10-	8-Aminoquinaldine	Crotonaldehyde diacetate	7	315
3,4-Dimethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	Glycerol	22	271
3,5-Dimethyl-1,10-	8-Amino-6-methylquinoline	$CH_2=C(CH_3)$ -	4	317
		CH(OCOCH ₃) ₂		
3,6-Dimethyl-1,10-	8-Amino-3,6-dimethylquinoline	Glycerol	3	317
3,7-Dimethyl-1,10-	8-Amino-4-methylquinoline	$CH_2 = C(CH_3)CH(OCOCH_3)_2$	2	315
3,8-Dimethyl-1,10-	8-Amino-3-methylquinoline	$CH_2=C(CH_3)CH(OCOCH_2)_2$	9	315
4,5-Dimethyl-1,10-	8-Amino-4,5-dimethylquinoline	Glycerol	24	317
4,6-Dimethyl-1,10-	8-Amino-4,6-dimethylquinoline	Glycerol	11	317
4,7-Dimethyl-1,10-	8-Amino-4-methylquinoline	CH3OCH2CH2C(OCH3)2CH3	7	315
5,6-Dimethyl-1,10-	8-Amino-5,6-dimethylquinoline	Glycerol	9	315
4,6-Diphenyl-1,10-	4,6-Diphenyl-8-aminoquinoline	Glycerol	10	282
4.7-Diphenyl-1,10-	4-Phenyl-8-aminoquinoline	C6H6COCH2CH2Cl	40	282
3,4,6-Trimethyl-1,10-	3,4,6-Trimethyl-8-aminoquinoline	Glycerol	19	271
3,4,7-Trimethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	CH ₂ COCH=CH ₂	31	271
3,4,8-Trimethyl-1,10-	3,4-Dimethyl-8-aminoquinoline	$CH_2 = C(CH_3)CH(OCOCH_3)_2$	9	271
3,5,6-Trihromo-1,10-	8-Amino-3,5,6-trihromoquinoline	Glycerol	27	316
3.5,6-Trimethyl-1,10-	8-Amino-5,6-dimethylquinoline	$CH_2 = C(CH_3)CH(OCOCH_3)_2$	9	315
3.5.7-Trimethyl-1,10-	8-Amino-4,6-dimethy lquinoline	$CH_2 = C(CH_3)CH(OCOCH_3)_2$	15	317
3.5.8-Trimethyl-1.10-	8-Amino-3,5-dimethylquinoline	$CH_2=C(CH_3)CH(OCOCH_3)_2$	9	317
3,6,7-Trimethyl-1,10-	8-Amino-4,5-dimethylquinoline	$CH_2=C(CH_3)CH(OCOCH_3)_2$	2	317
4.5.7-Trimethyl-1.10-	8-Amino-4,6-dimethylquinoline	CH3COCH=CH2	1	317
3,5,6,8-Tetrahromo-1,10-	8-Amino-3,5,6-trihromogninoline	$CH_2 = C(B_r)CH(OCOCH_2)_2$	4	316
3,4,6,7-Tetramethyl-1,10-	3,4,6-Trimethyl-8-aminoquinoline	CH3COCH=CH2	5	271
3.4.7.8 Table 1.10-	3.4.6-Trimethyl-S-aminoquinoline	CH_COOK(OH_OCH_OU	9	271

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

CH2COCH(CH2)CH2OH

CH2=C(CH2)CH(OCOC112)2

3,4,7,8-Tetramethyl-1,10- 3,4-Dimethyl-S-aminoquinoline

3.5,6,8-Tetramethyl-1,10- 8-Amino-3,5,6-trimethylquinoline

20

22

271

315

TABLE V-Continued

PHENANTHROLINES

B. 1.7-Phenanthrolines





6-Bromo-1.7-5-Nitro-1,7-2-Hydroxy-1,7-6-Hydroxy-1,7-8-Hydroxy-1,7-10-Hydroxy-1,7-2-Methyl-1,7-(together with the linear isomer 2-methyl-1,9-anthrazoline) 6-Methyl-1.7-2-Hydroxy-4-methyl-1.7-

Reactants			Relec-
Amine	Second Component	Yield %	ences
m-Phenylenediamine	Glycerol	80	36, 43, 70 71, 72 183
4-Bromo-m-phenylenediamine	Glycerol	30	131
5-Nitro-m-phenylenediamine	Glycerol	44	222
2-Hydroxy-7-aminoquinoline	Glycerol	50	185
5-Amino-8-hydroxyquinoline	Glycerol	40	131
2,4-Dinitrophenol	Glycerol	10	255
2-Hydroxy-5-aminoquinoline	Glycerol	62	165
4-Hydroxy-5-aminoquinoline	Glycerol	60	240
2-Methyl-7-aminoquinoline	Glycerol	_	163
8-Methyl-5-aminoquinoline	Glycerol	_	280
2-Hydroxy-4-methyl-7-aminoquinoline	Glycerol	60	185
4-Hydroxy-2-methyl-5-aminoquinoline	Glycerol	60	185

10-Hydroxy-8-methyl-1,7-Phenanthroline



10/	N.
8	T
8 7 L	6_6
N	•

(20)	'N
la , 1 5	
N 6	
4,7-Phenanthrolis	ne

1,2,3,4-Tetrahydro-4,7-
or the linear isomer
1,2,3,4-tetrahydro-
1,6-anthrazoline
6-Bromo-4,7-
1-Hydroxy-4,7-
3-Hydroxy-4,7-
1-Hydroxy-3-methyl-4,7-
3-Hydroxy-1-methyl-4,7-
3-Keto-4-methyl-4,7-

1,8-Phenanthroline
5-Methyl-1,6-phenanthroli
5-Methyl-1,6-anthrazoline
2-Hydroxy-4,5,10-tri-
methyl-1,6-anthrazoline
Mates D.C

1,3-Dimethyl-4,7-3,8-Dimethyl-4,7-5,6-Benzo-4,7-

Reactants

Second Component

Glycerol

Yield

%

60

Refer-

ences

36, 70, 71, 79

C. 4.7-Phenanthrolines

Amine

p-Phenylenediamine

				12
	p-Nitroaniline	Glycerol	46	73
	6-Aminoquinoline	Glycerol	100	50,73,180
	1,2,3,4-Tetrahydro-6-aminoquinoline	Glycerol	_	217
	0.7			
	8-Bromo-6-aminoquinoline	Glycerol	60	131
	4-Hydroxy-6-aminoquinoline	Glycerol	Good	186
_	2-Hydroxy-6-aminoquinoline	Glyecrol	Quant.	74
7-	4-Hydroxy-2-methyl-6-aminoquindine	Glycerol	88	186
7-	2-Hydroxy-4-methyl-6-aminoquinoline	Glyrerol	88	50, 186
	2-Keto-1-methyl-6-aminoquinoline	Glycerol	55	74
	2,4-Dimethyl-6-aminoquinoline	Glycerol		143
	p-Phenylenediamine	CH2CH=CHCHO		244
	1,4-Diaminonaphthalene	Glycerol		143
	D. Other Phenanthrolines			
	5-Aminoisoquinoline	Glycerol	5	75
roline	2-Methyl-4-aminoquinoline	Glycerol	6	81, 82
line	5-Methyl-6-acetylaminoquinoline	Glycerol	_	50
line	2-Hydroxy-4,5,8-trimethyl-6-amino- quinoline	Glycerol	_	50

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

TABLE VI

MISCELLANEOUS QUINOLINES

Reactants

				
		Second	Yiel	d Refer-
Product	Amine	Component	%	ences
5.6-Trimethylenequinoline	3,4-Trimethyleneaniline	Glycerol	6	94
6,7-Trimethylenequinoline	3,4-Trimethyleneaniline	Glycerol	54	94
7,8-Trimethylenequinoline	2,3-Trimethyleneaniline	Glycerol	60	147
7,12-Diketonaphtho(2,3-h)quinoline	1-Amino-9,10-diketoanthracene	Glycerol	_	43, 60, 61
5,6-Dihydroxy-7,12-diketonaphtho- (2,3-h)quinoline	1-Amino-3,4-dihydroxy-9,10-diketo- anthracene	Glycerol	_	43
8-Amino-9-methyl-7,12-diketonaph- tho(2,3-h)quinoline	1,5-Diamino-2-methyl-9,10-diketo- anthracene	Glycerol	97	63
10-Methyl-11-amino-7,12-diketonaph- tho(3,2-h)quinoline	1,8-Diamino-2-methyl-9,10-diketo- anthracene	Glycerol	_	63
Naphtho(2,3-f)quinoline	2-Aminoanthracene	Glycerol	_	64, 164
7,12-Diketonaphtho(2,3-f)quinoline	2-Amino-9,10-diketoanthracene	Glycerol	_	62
3-Methyl-7,12-diketonaphtho(2,3-f)-quinoline	2-Amino-9,10-diketoanthracene	Paraldehyde		60
5,6-Dihydroxy-7,12-diketonaphtho- (2,3-f)quinoline	2-Amino-3,4-dihydroxy-9,10-diketo- anthracene	Glycerol		43, 59, 64
6,7-Benz-12-ketonaphtho(2,3-f)- quinoline	2-Amino-9,10-diketoanthracene	Glycerol	-	171
Naphtho(1,2-h)quinoline	1-Aminophenanthrene	Glycerol	_	55
Naphtho(2,1-f)quinoline	2-Aminophenanthrene	Glycerol	90	56
5,6-Dihydronaphtho(1,2-q)quinoline	2-Amino-9.10-dihydrophenanthrene	Glycerol	50	56
Naphtho(1,2-f)quinoline	3-Aminophenanthrene	Glycerol	45	56
Naphtho(2,1-h)quinoline	4-Aminophenanthrene	Glycerol	20	55
Dibenzo(f,h)quinoline	9-Aminophenanthrene	Glycerol	60	54
Pyrenoline	3-Aminopyrene	Glycerol	_	57
11-Indeno(2,1-f)quinoline	2-Aminofluorene	Glycerol	_	65
1,5-Naphthyridine	3-Aminopyridine	Glycerol	28	66, 274,
-,,,,	o managa managa			513
2-Hydroxy-1,5-naphthyridine	3-Amino-6-hydroxypyridine	Glycerol	15	66, 314
Thieno(2,3-b)pyridine	2-Aminothiophene	Glycerol	5	67
2-Keto-1,2-dihydro-1-oxa-8-aza- phenanthrene	6-Aminocoumarin	Glycerol	57	68, <i>300</i>
9-Methyl-2-keto-1,2-dihydro-1-oxa- 8-azaphenanthrene	6-Nitro-7-methylcoumarin	Glycerol	35	68 .
4.9-Dimethyl-2-keto-1,2-dihydro- 1-oxa-8-azaphenanthrene	6-Nitro-4,7-dimethylcoumarin	Glycerol	20	68
9,10-Benz-2-keto-1,2-dihydro-1-oxa- 8-azaphenanthrene	6-Nitro-1,2-α-naphthapyrone	Glyeerol	30	68
4-Methyl-9,10-benz-2-keto-1,2-dihy- dro-1-oxa-8-azaphenanthrenc	6-Nitro-4-methyl-1,2-α-naphtha- pyrone	Glycerol	50	68
Benzofuro(2,3-f)quinoline	3-Aminodibenzofuran	Glycerol		95, 96, 97
Benzofuro(3,2-g)quinoline	3-Aminodibenzofuran	Glycerol		<i>95</i> , 96, 97
5-Nitrobenzofuro(2,3-f)quinoline	3-Amino-2-nitrodibenzofuran	Glycerol	24	97
Benzofuro(3,2-f)quinoline	2-Aminodibenzofuran	Glycerol	_	96
Benzoluro(2,3-g)quinoline	2-Aminodibenzofuran	Glycerol		96
5-Benzenesulfonamidobenzofuro- (3,2-f)quinoline	2-Amino-3-benzenesulfonamido- dibenzofuran	Glycerol	45	97
12-Xanthono(2,1-b) pyridine	2-Aminoxanthone	Glycerol	_	69
10-Nitro-12-xanthono(2,1-b)pyridine	2,7-Dinitroxanthone	Glycerol	_	69
Pyridino(2',3',4,5)benzothiazole	4-Aminobenzothiazole	Glycerol	30	305
Pyridino(2',3',6,7)benzothiazole	6-Aminobenzothiazole	Glycerol	50	304
2-Methylpyridino(3',2',4,5)benzo- thiazole	5-Amino-2-methylbenzothiazole	Glycerol	_	302

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

TABLE VI—Continued MISCELLANEOUS QUINOLINES

Reactants

Product 3-Phenyl-3-triazolobenzo(f)quinoline 2-Phenyl-2-triazolobenzo(f)quinoline 2-p-Tolyl-2-triazolobenzo(f)quinoline or	Amine 1-Phenyl-5-amino-1-benzotriazole 2-Phenyl-5-amino-2-benzotriazole 2-p-Tolyl-5-nitro-2-benzotriazole	Second Component Glycerol Glycerol Glycerol	Yield % — —	References 322 322 137
2-p-Tolyl-2-triazolobenzo(p)quinoline Pyridino(3',2',4,5)-benzothiadiazole 3-Phenyl-3-imidazo(f)quinoline 2-Phenyl-3-imidazo(f)quinoline 3-p-Tolyl-3-imidazo(f)quinoline 1-Phenyl-4-chloro-1-imidazo(g)quinoline	5-Aminobenzothiadiazole 1-Phenyl-5-aminobenzimidazole 2-Phenyl-5-aminobenzimidazole 1-p-Tolyl-5-aminobenzimidazole 1-Phenyl-4-chloro-5-aminobenz- imidazole	Glycerol Glycerol Glycerol Glycerol	- 35 -	303 322 322 322 322 322
nne 1-p-Tolyl-4-chloro-1-imidazo(g)quino- line 2-Phenyl-4-bromo-1-imidazo(g)quino- line	1-p-Tolyl-4-chloro-5-aminobenz- imidazole 2-Phenyl-4-hromo-5-aminobenz- imidazole	Glycerol Glycerol	_	322 322
1-Pyrazolo(3,4-f)quinoline 9-Chloro-1-pyrazolo(4,3-g)quinoline Quinolino(8,7-h)quinoline 9,16-Diketodipyridoanthracene 9,10-Diketodipyridoanthracene 9,10-Diketodipyridoanthracene Dimethyldipyridoacridine Dipyrido(2,3-f/h)quinoline Di-6-quinolyl oxide	6-Aminoindazole 6-Amino-7-chloroindazole 1,5-Diamino-9,10-anthraquinone 2,6-Diamino-9,10-anthraquinone 2,7-Diamino-9,10-anthraquinone 3,6-Diamino-9,10-anthraquinone 3,6-Diaminoacridine 1,3,5-Triaminobenzene 4,4'-Diaminodiphenyl oxide	Glycerol Glycerol Glycerol Glycerol Glycerol Glycerol CH ₂ CH=CHCHO Glycerol Glycerol	30	322 322 194 61 62 62 244 210 310

Note: References 116-322 are listed on pp. 94-98. Where one reference is italicized, the yield reported is taken from that reference.

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CHAPTER 3

CARBON-CARBON ALKYLATIONS WITH AMINES AND AMMONIUM SALTS

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INTRODUCTION

This chapter is a review of those reactions of compounds containing labile amino groups in which a carbon-carbon bond is formed by amine replacement, as, for example, in the alkylation of diethyl malonate by 1-dimethylamino-3-butanone.

 $CH_3COCH_2CH_2N(CH_3)_2 + CH_2(CO_2C_2H_5)_2 \rightarrow$

$$CH_3COCH_2CH_2CH(CO_2C_2H_5)_2 + (CH_3)_2NH$$

In the most general terms, these alkylation reactions may be written

$$ZCH_2NR_1R_2 + HY \rightarrow ZCH_2Y + HNR_1R_2$$

OL

where

$$ZCH_2\overset{+}{N}R_1R_2R_3X^- + MY \rightarrow ZCH_2Y + NR_1R_2R_3 + MX$$

$$Z = R - C - CH -$$
, H , $CH -$, etc.;

HY = hydrogen cyanide, active methyl and methylene compounds; and MY = alkali cyanides, sodio derivatives of active methyl or methylene compounds, Grignard reagents, or organolithium compounds.

Attention has been given primarily to reactions of amines that can be prepared by the Mannich reaction (Mannich bases), but, for com-

¹ Blicke in Adams, O-panic Reactions, Vol. I, p. 303, John Wiley & Sons, 1942.

parison, analogous reactions of simpler quaternary ammonium salts have been included in the discussion and tables. A number of reactions which are closely related to these simple alkylations but follow a somewhat different pattern are discussed in the Related Reactions section and are not included in the tables.

SCOPE AND LIMITATIONS

General Considerations

The most important groups of compounds capable of engaging in carbon-carbon alkylations by amine replacement are:

- (a) Simple quaternary ammonium salts containing benzyl and methyl radicals. The general formulation of carbon-carbon alkylation with such salts corresponds to the third equation on p. 102
- (b) Tertiary amines that can be prepared from ketones, phenols, heterocyclic compounds, and nitro compounds by the Mannich reaction.¹

$$\begin{array}{c|c} O & H \\ \hline R-C-C-H+CH_2O+HNR_2\cdot HCI \rightarrow \\ \hline R-C-C-C+CH_2NR_2\cdot HCI+H_2O \\ \hline R' \\ \end{array}$$

 $\mathrm{CH_3CH_2CH_2NO_2} + \mathrm{CH_2O} + \mathrm{HNR_2} \, \rightarrow \, \mathrm{CH_3CH_2CH(NO_2)CH_2NR_2} + \mathrm{H_2O}$

The general form of the reactions of carbon-carbon alkylations by amine replacement undergone by these Mannich bases is shown in the second equation on p. 102.

(c) Quaternary salts of Mannich bases, which can be formed by reaction of the tertiary amines with alkyl halides or dimethyl sulfate.

$$ZCH_2NR_2 + R'X \rightarrow ZCH_2NR_2R'X^-$$

The general form of the reactions undergone by these salts is shown in the third equation on p. 102.

Structural Considerations

Structure of the Alkylating Radical. The ability to form a conjugated unsaturated system by amine elimination seems to be the main structural requirement for facile carbon-carbon alkylations by amine replacement with tertiary amines (see p. 126). The structural features required for amine elimination are indicated in formulas I and II. An enolizable hydrogen atom must be so located that when it and the dialkylamino

group are removed from the molecule a conjugated unsaturated system can be established by electron transfer.

The structural characteristics necessary for easy carbon-carbon alkylations with quaternary ammonium salts are similar. A number of quaternary salts that cannot undergo amine elimination can be used as alkylating agents, although in general the reactions are much slower than those of quaternary salts which can suffer amine elimination. Where amine climination is not possible, the structural requirement of the alkylating radical appears to be either the presence of an allylic system, as in benzyl, 1-methylskatyl (III), and furfuryl radicals, or freedom from steric hindrance to rearward attack, as in the methyl radical.

Structure of the Amino Group Replaced. The structure of the amino group replaced in carbon-carbon alkylations of this type is of some importance in the economic and operational aspects of these reactions. The presence of certain amino groups that could undergo alkylation by the alkylating radical, such as derivatives of aniline, is probably undesirable in some of these reactions.

Structure of the Substance To Be Alkylated. Only those substances that can easily form anions can be alkylated by Mannich bases or quaternary salts. Active methylene compounds and their sodio derivatives, hydrogen cyanide and its salts, and organometallic compounds such as Grignard reagents and alkyl- or aryl-lithium compounds constitute the principal members of this class of substances.

The carbon-carbon alkylations with amines and ammonium salts to be considered in detail are the following.

- (a) Replacement of amino groups by cyanide
- (b) Alkylation of active methyl and methylene compounds
 - 1. Alkylation of aliphatic nitro compounds
 - 2. Alkylation of ketones and β -keto esters
 - 3. Alkylation of esters
 - 4. An alkylation of indole
- (c) Amine replacement reactions of quaternary salts with organometallic compounds.

Replacement of Amino Groups by Cyanide

Quaternary Ammonium Salts and Alkali Cyanides. Quaternary ammonium cyanides are difficult to prepare, but mixtures of certain quaternary ammonium salts with alkali cyanides decompose when strongly heated in a manner expected of quaternary ammonium cyanides. The reactions are analogous to those of quaternary ammonium halides in that benzyl and methyl groups are cleaved from the quaternary nitrogen atom and couple with the anion of the salt. In at least one reaction, however, olefin formation, similar to that found in the Hofmann exhaustive methylation, occurs more readily than does simple amine replacement.²

When tetramethylammonium cyanide is heated, acetonitrile, methylcarbylamine, and trimethylamine are formed.³ Acetonitrile and methylethylaniline are formed when a mixture of potassium cyanide and dimethylethylanilinium iodide is distilled to dryness.⁴

² Snyder and Brewster, J. Am. Chem. Soc., 71, 291 (1949).

³ Thompson, Ber., 16, 2338 (1883).

⁴ von Meyer and Schwabe, Abhandl. math.-phys. Klasse sächs. Ges. Wiss., 31, 179 (1908) [Chem. Zentr., 80, II, 1800 (1909); C. A., 5, 887 (1911)].

Although benzyldimethylanilinium halides do not react appreciably with sodium cyanide in boiling water, benzyl cyanide is formed when an aqueous solution of the two salts is distilled to dryness. Similarly, the methiodide of 1-dimethylaminomethyl-2-methoxynaphthalene (IV, $R = CH_3$) reacts with sodium cyanide to form 2-methoxy-1-naphthylacetonitrile (V, $R = CH_3$) only when an aqueous solution of the two salts is evaporated to dryness and distilled in vacuum at temperatures above 150°. On the other hand, when a mixture of sodium cyanide and N,N,N-trimethyl- α -phenylethylammonium iodide (VI) was similarly

$$\begin{array}{c} CH_2N(CH_3)_2 \\ \hline \\ OR \\ \hline \\ V \end{array} \begin{array}{c} CH_2CN \\ \hline \\ V \end{array}$$

treated, styrene was formed and no hydratroponitrile could be detected in the reaction products.²

Although none of the reactions described above is of preparative interest, since the corresponding methyl and benzyl halides are readily available, the analogous reactions of the quaternary salts of Mannich bases derived from indole are useful in the preparation of indoleacetonitriles. The methiodide of gramine ^{6a,b,c} (3-dimethylaminomethylindole, VIIa) reacts with potassium silver cyanide in boiling water to form indole-3-acetonitrile (VIII), isolated as the acid in 46% yield. ^{6c,7} The methosulfate of gramine reacts readily with potassium cyanide in aqueous ethanol to form the same nitrile (VIII) (isolated as the acid in 50% yield from gramine). ^{8,8a} The quaternary salt of gramine is formed

⁵ Snyder and Speck, J. Am. Chem. Soc., 61, 668 (1939).

⁶ Snyder and Brewster, J. Am. Chem. Soc., 71, 1058 (1949).

Schramm, J. Am. Chem. Soc., 73, 2961 (1951).
 Schöpf and Thesing, Angew. Chem., 63, 377 (1951).

⁶c Geissman and Armen, J. Am. Chem. Soc., 74, 3916 (1952).

⁷ Snyder, Smith, and Stewart, J. Am. Chem. Soc., 66, 200 (1944).

⁸ Heidelberger, J. Biol. Chem., 179, 139 (1949).

⁸a Thesing and Schülde, Chem. Ber., 85, 324 (1952).

in situ by the addition of dimethyl sulfate to the solution of gramine and potassium cyanide. The methiodide of 1-methylgramine (IX) reacts with hot aqueous sodium cyanide to give mainly the expected product, 1-methyl-3-indoleacetonitrile (X, 60-64%), together with smaller amounts of 1,3-dimethyl-2-cyanoindole (XI, 4%), apparently by an allylic rearrangement during the alkylation process. The Mannich bases of N-methyl- and N-phenyl-pyrrole yield the normal products only. 3c

$$CH_{2}\overset{+}{\text{N}}(CH_{3})_{3}\text{ I} \xrightarrow{\text{NaCN}}$$

$$CH_{3}\overset{+}{\text{IX}}$$

$$CH_{2}\text{CH}_{2}\text{CN} + \text{CH}_{3}\overset{+}{\text{CN}} + \text{N(CH}_{3})_{3} + \text{NaI}$$

$$CH_{3}\overset{+}{\text{CH}_{3}}$$

$$CH_{3}\overset{+}{\text{CH}_{3}}$$

$$CH_{3}\overset{+}{\text{NaI}}$$

In a similar fashion, furfuryltrimethylammonium iodide (XII, R = H) yields a mixture of furfuryl cyanide (XIII, R = H, 27%) and 2-cyano-5-methylfuran (XIV, 5%), and 5-methylfurfuryltrimethylammonium iodide (XII, $R = CH_3$) gives 5-methylfurfuryl cyanide (XIII, $R = CH_3$) in 37% yield.¹⁰

The methiodide of β -dimethylaminopivalophenone (XV) reacts with sodium cyanide when an aqueous solution of the two salts is distilled to form β -dimethylaminopivalophenone (XV) and, presumably, acetonitrile.¹¹

⁹ Snyder and Eliel, J. Am. Chem. Soc., 70, 1703, 1857 (1948).

⁹a Herz and Rogers, J. Am. Chem. Soc., 73, 4921 (1951).

Eliel and Peckham, J. Am. Chem. Soc., 72, 1209 (1950).
 Snyder and Brewster, J. Am. Chem. Soc., 71, 1061 (1949).

Tertiary Amines and Hydrogen Cyanide. Tertiary amines capable of eliminating a secondary amine to form a conjugated unsaturated structure can react with hydrogen cyanide to form nitriles by amine replacement.

3-Dialkylaminomethylindoles (VII) react with hydrogen cyanide in benzene solution at 150° to form indole-3-acetonitrile (VIII); 12 under similar conditions 1-dimethylaminomethyl-2-hydroxynaphthalene (IV, R = H) reacts with hydrogen cyanide to form 2-hydroxy-1-naphthaleneacetonitrile (V, R = H).12 No information on the yields obtainable by this process is available.

Hydrochlorides of a number of ketonic Mannich bases have been found to react readily with alkali metal cyanides in hot water to form γ -ketonitriles in good yield. No successful application of this reaction to wholly aliphatic ketonic Mannich bases has been reported; the hydrochloride of 2-dimethylaminomethylcyclohexanone (XVI) formed only a resin or oil when heated with potassium cyanide in aqueous solution.13 Ketonic Mannich base hydrochlorides of structure XVII have been found to react satisfactorily with aqueous potassium cyanide when R is furyl, benzofuryl, thienyl, phenyl, 3-hydroxy- and 3-methoxyphenyl, 4-methyl-, 4-chloro-, 4-bromo-, 4-hydroxy-, and 4-methoxy-

$$\begin{array}{c} R_1 \\ \downarrow \\ RCOCCH_2N(CH_3)_2 \cdot HCl + KCN \rightarrow RCOCCH_2CN + HN(CH_3)_2 + KCl \\ \downarrow \\ R_2 \\ XVII \quad R_1 = R_2 = H \end{array}$$

phenyl; 3,4-dimethoxyphenyl, α - or β -naphthyl. The hydrochloride of β -dimethylamino-3-nitropropiophenone formed resins when heated with aqueous potassium cyanide.13

Substituents on the carbon atom adjacent to the carbonyl group appear to interfere with the reaction with cyanides. The hydrochloride of α -dimethylaminomethylpropiophenone (XVII, $R_1 = H$, $R_2 = CH_3$) formed a resin or oil,13 and the hydrochloride of dimethylaminopivalophenone (XVII, R₁ = R₂ = CH₃) underwent a reverse Mannich reaction to form isobutyrophenone.11

¹² Salzer and Andersag, U. S. pat. 2,315,661 [C. A., 37, 5418 (1943)]; U. S. PB 706, Dept. of Commerce, Washington, D. C.

¹³ Knott, J. Chem. Soc., 1947, 1190.

It has been reported that the salts of Mannich bases made from piperidine or morpholine do not react under conditions ¹³ suitable for dimethylamine derivatives. It seems likely that this is at least partly due to the fact that the amines being replaced are less volatile than the solvent.

Tertiary Amines and Alkali Cyanides. The Mannich bases of phenols and indoles react with sodium cyanide in hot aqueous ethanol to form sodium salts of aryl- and indole-acetic acids.¹² Little information on yields and the by-products formed is available, though it is reported that condensation products are formed from phenolic Mannich bases. This is not surprising since phenolic Mannich bases readily undergo self-alkylation in weakly alkaline solution to form diarylmethanes.¹⁴

$$2ZCH_2NR_1R_2 + H_2O \rightarrow ZCH_2Z + CH_2O + 2HNR_1R_2$$

In the reaction of 1-dimethylaminomethyl-2-naphthol with sodium cyanide it was found that 2-hydroxy-1-naphthaleneacetic acid (XVIII) could be isolated in 47% yield, and the diarylmethane (XIX) was formed in at least 20% yield. It seems likely that diarylmethane formation would be a major side reaction in any similar application of this method and that phenolic Mannich bases containing unsubstituted ortho or para positions would form appreciable amounts of polymeric materials, as has

$$\begin{array}{c} \text{CH}_2\text{N}(\text{CH}_3)_2 & \xrightarrow{\text{NaCN, H}_2\text{O}} & \text{CH}_2\text{CO}_2\text{H} & + \\ \text{OH} & & \text{XVIII (47\%)} & \\ \end{array}$$

been observed in the reaction of 6-dimethylaminomethylguaiacol with sodium cyanide. 15a

Several 3-dialkylaminomethylindoles (VII) have been subjected to reaction with cyanide, but information as to yields is available only for dimethylaminomethylindole (gramine, VIIa) which in hot aqueous

¹⁴ Auwers and Dombrowski, Ann., 344, 280 (1906).

¹⁵ J. Brewster, doctoral thesis, University of Illinois, Urbana, Ill., 1948.

¹⁵a Eliel, J. Am. Chem. Soc., 73, 43 (1951).

 R_2

XVII $R_1 = R_2 = H$

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$$\begin{array}{c} \text{CH}_2\text{N}(\text{CH}_3)_2 \\ \text{XVI} \\ \\ \text{R}_1 \\ \\ \text{RCOCCH}_2\text{N}(\text{CH}_3)_2 \cdot \text{HCl} + \text{KCN} \rightarrow \text{RCOCCH}_2\text{CN} + \text{HN}(\text{CH}_3)_2 + \text{KCl} \end{array}$$

phenyl; 3,4-dimethoxyphenyl, α - or β -naphthyl. The hydrochloride of β -dimethylamino-3-nitropropiophenone formed resins when heated with aqueous potassium eyanide.¹³

R.

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Salzer and Andersag, U. S. pat. 2,315,661 [C. A., 37, 5418 (1943)]; U. S. PB 706,
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$$CH_2$$
 + $HN(CH_3)_2$ + NH_3 + CH_2O
 $XIX (20\%)$

been observed in the reaction of 6-dimethylaminomethylguaiacol with sodium cyanide. 15a

Several 3-dialkylaminomethylindoles (VII) have been subjected to reaction with cyanide, but information as to yields is available only for dimethylaminomethylindole (gramine, VIIa) which in hot aqueous

¹⁴ Auwers and Dombrowski, Ann., 344, 280 (1906).

¹³ J. Brewster, doctoral thesis, University of Illinois, Urbana, Ill., 1948.

¹⁵¹ Eliel, J. Am. Chem. Soc., 73, 43 (1951).

ethanol gave a 69% yield of 3-indoleacetic acid (XX) and a 20% yield of 3-indoleacetamide with little or no diindolylmethane. Indoleacetamide may be hydrolyzed to the acid in good yield.

$$\mathrm{CH_{2}CO_{2}H}$$
 $\mathrm{CH_{2}N(CH_{3})_{2}}$
 $\mathrm{CH_{3}}$
 $\mathrm{CH_{3}}$
 XXI

Compounds that cannot suffer amine elimination, such as 1-methylgramine 17 (XXI) and 1-dimethylaminomethyl-2-methoxynaphthalene 6 (IV, R = CH₃) fail to react with sodium cyanide under the above conditions.

Alkylation of Active Methyl and Methylene Compounds

Alkylation of Aliphatic Nitro Compounds. Alkylations of aliphatic nitro compounds by *p*-nitrobenzyltrimethylammonium iodide and Mannich bases of indole, of ketones, and of aliphatic nitro compounds have been reported.

Gramine (VIIa) reacts smoothly with 1- or 2-nitropropane in the presence of sodium hydroxide to give good yields of monoalkylated nitro compound; much lower yields are obtained with nitroethane. Only

diskatylnitromethane (XXII) was obtained by alkylation of nitromethane under these conditions 18

¹⁶ Snyder and Pilgrim, J. Am. Chem. Soc., 70, 3770 (1948).

Snyder and Eliel, J. Am. Chem. Soc., 71, 663 (1949).
 Snyder and Katz, J. Am. Chem. Soc., 69, 3140 (1947).

$$2 \xrightarrow{\text{CH}_2\text{N}(\text{CH}_3)_2} + \text{CH}_3\text{NO}_2 \xrightarrow{\text{NaOH}}$$

$$\text{H}_{\text{VII}a}$$

$$\text{CH}_2\text{CH}_2\text{CHNO}_2 + 2(\text{CH}_3)_2\text{NH}$$

$$\text{H}_{\text{VII}}$$

Ethyl nitroacetate is dialkylated with gramine in the presence of ethanol and sodium ethoxide ¹⁸ or in the presence of powdered sodium hydroxide in xylene. ¹⁹ Skatylnitroacetic ester (XXIII), which can be converted to tryptophan in good yield, is obtained from gramine and ethyl nitroacetate in xylene solution in the absence of any catalyst; ¹⁹ diethyl nitromalonate may also be alkylated by means of gramine and the product may be converted to tryptophan. ²⁰

Ketonic Mannich bases react rapidly with nitromethane in the presence of alkaline catalysts, as sodium methoxide or ethanolic potassium hydroxide, to form mono-, di-, or tri-alkylated nitromethanes. Thus, with the Mannich bases of acetone (XXIV), cyclohexanone (XVI), acetophenone (XXV), and 4-methoxy- and 3,4-dimethoxy-acetophenone, monoalkylated products are formed from nitromethane in the presence of sodium ethoxide. Some dialkylated product is formed from the

RCOCHCH₂N(CH₃)₂ + CH₃NO₂
$$\xrightarrow{\text{Base}}$$

R'

RCOCHCH₂CH₂NO₂, (RCOCHCH₂)₂CHNO₂, or (RCOCHCH₂)₃CNO₂

R'

R'

R'

¹⁹ Lyttle and Weisblat, J. Am. Chem. Soc., 69, 2118 (1947); Weisblat and Lyttle, U. S. pat. 2,557,041 [C. A., 46, 1593 (1952)].

²⁹ Weisblat and Lyttle, J. Am. Chem. Soc., 71, 3079 (1949); U. S. pat. 2,528,928 [C. A., 45, 3870g (1951)].

²¹ Reichert and Posemann, Arch. Pharm., 275, 67 (1937).

Mannich base of 3,4-dimethoxyacetophenone. Di- and tri-alkylated nitromethanes are formed by reaction of the Mannich base of acetophenone, nitromethane, and ethanolic potassium hydroxide.

$$\begin{array}{ccc} CH_3COCH_2CH_2NR_2 & C_6H_5COCH_2CH_2NR_2 \\ xxiv & xxv \end{array}$$

1- and 2-Nitropropane can be alkylated by the Mannich base derived from 1-nitropropane. 21c, b The reaction fails with the Mannich base of 2-nitropropane.

Alkylation of Ketones and β-Keto Esters. Many alkylations of ketones and β-keto esters by means of Mannich bases have been reported.21c The principal interest in these reactions has been in the prepa-

21a Snyder and Hamlin, J. Am. Chem. Soc., 72, 5082 (1950).

Other examples are reported by Gill, James, Lions, and Potts, J. Am. Chem. Soc., 74, 4923 (1952).

the For more recent examples see Barltrop and Saxton, J. Chem. Soc., 1952, 1038; Gunstone and Heggie, ibid., 1952, 1437.

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$$\begin{array}{ccc} \mathrm{CH_{3}COCH_{2}CH_{2}NR_{2}} & & \mathrm{C_{6}H_{5}COCH_{2}CH_{2}NR_{2}} \\ & & \mathrm{xxv} \end{array}$$

1- and 2-Nitropropane can be alkylated by the Mannich base derived from 1-nitropropane. The reaction fails with the Mannich base of 2-nitropropane.

Alkylation of Ketones and β -Keto Esters. Many alkylations of ketones and β -keto esters by means of Mannich bases have been reported.^{21c} The principal interest in these reactions has been in the preparation.

Snyder and Hamlin, J. Am. Chem. Soc., 72, 5082 (1950).
 Other examples are reported by Gill, James, Lions, and Potts, J. Am. Chem. Soc., 4323 (1952).

ne For more recent examples see Barltrop and Saxton, J. Chem. Soc., 1952, 1038; Gunstone and Heagle, Eds., 1952, 1437.

$$\begin{array}{c} \text{CH}_3\text{COCH}_2\text{CH}_2\overset{\dagger}{\text{N}}(\text{CH}_3)(\text{C}_2\text{H}_5)_2\text{ I} \\ \\ \text{H}_3\text{C} \\ \\ \text{O} \end{array} \qquad \begin{array}{c} \text{CH}_3\text{COCH}_2\text{CH}_2\overset{\dagger}{\text{CH}_3} \\ \\ \text{CH}_3\text{COCH}_2\text{CH}_2\overset{\dagger}{\text{N}}(\text{CH}_3)(\text{C}_2\text{H}_5)_2\text{ I} \\ \\ \text{H} \\ \\ \text{O} \end{array} \qquad \begin{array}{c} \text{CH}_3\text{COCH}_2\text{CH}_2 \\ \\ \text{CH}_3\text{COCH}_2\text{CH}_2 \\ \\ \text{O} \end{array} \qquad \begin{array}{c} \text{CH}_3 \\ \\ \text{O} \end{array} \qquad \begin{array}{c} \text{CH}_3 \\ \\ \text{CH}_3\text{COCH}_2\text{CH}_2 \\ \\ \text{O} \end{array} \qquad \begin{array}{c} \text{OCH}_3 \\ \\ \text{OCH}_3 \\ \\ \text{OCH}_3 \end{array} \qquad \begin{array}{c} \text{OCH}_3 \\ \\ \text{OCH}_3 \\ \\ \text{OCH}_3 \end{array} \qquad \begin{array}{c} \text{OCH}_3 \\ \\ \text{OCH}_3$$

It will be noted that the alkylation products of ketones or β -keto esters with a ketonic Mannich base are δ-diketones, many of which can form cyclohexenone derivatives by internal aldol condensation as in the examples cited above. Often, as above, such cyclizations occur during alkylation. These reactions may be used to form simple cyclohexenone derivatives, such as the terpenes carvenone (XXIX) and piperitone 32, 32a, 32b (XXX), bicyclic terpenes containing angular methyl groups

$$H_3C$$
 $CH(CH_3)_2$
 H_3C
 $CH(CH_3)_2$
 $CH(CH_3)_2$

such as the cyperones 33 (XXXI), polynuclear aromatic hydrocarbons, 25 fused ring systems related to the steroids and containing angular methyl groups,34,35 compounds related to alkaloids and containing angular

Downes, Gill, and Lions, Australian J. Sci., 10, 147 (1948) [C. A., 42, 7257 (1948)].

²⁴ Downes, Gill, and Lions, J. Am. Chem. Soc., 72, 3464 (1950).

²⁵ Gill and Lions, J. Am. Chem. Soc., 72, 3468 (1950).

³ Adamson, McQuillin, Robinson, and Simonsen, J. Chem. Soc., 1937, 1576; McQuillin, ibid., 1951, 716.

³⁴ Martin and Robinson, J. Chem. Soc., 1943, 491; 1949, 1866.

Sec., 1949, 1855.

used as the base. Only a few alkylations of a ketone by a free ketonic Mannich base (tertiary amine) have been reported. One is the alkylation of 2-phenylcyclohexanone (XXVIII) with a Mannich base of acetone (XXIV), in the presence of one equivalent of sodium amide, which proceeds in 42% yield.³⁰ In two other cases, the bases were employed as hydrochlorides with sodium hydroxide or potassium t-butoxide as catalyst.

The yield of alkylation product may be increased by formylating the ketone first by means of methyl formate. The resulting α -hydroxymethyleneketone (which is considerably more acidic than the parent ketone) is then alkylated in good yield with the methiodide of the ketonic Mannich base in the presence of sodium methoxide, and the hydroxymethylene group is finally removed by basic cleavage at the same time cyclization is effected.^{30a, b}

more readily alkylated than active methylene groups. An active methylene group bearing a phenyl group is more readily alkylated than one bearing only alkyl groups. The following examples illustrate these principles.^{25,31}

¹³ Bockelheide, J. Am. Chem. Soc., 69, 790 (1947).

et al., J. Am. Chem. Soc., 72, 2388 (1950); see, however, Woodward

²²⁵ Wilds and Werth, J. Org. Chem., 17, 1149, 1154 (1952).
²¹ Crowley and Robinson, J. Chem. Soc., 1938, 2001.

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$$\begin{array}{c|c} R & \xrightarrow{HCO_2CH_3} & \\ \hline & NaOCH_3 & \\ \end{array} \qquad \begin{array}{c} CHOH \\ \end{array} \qquad + \\ \end{array}$$

$$\text{CH}_3\text{COCH}_2\text{CH}_2\text{N}(\text{CH}_3)(\text{C}_2\text{H}_5)_2\text{ I}^-\xrightarrow{\text{NaOCH}_8}\text{R}\xrightarrow{\text{CH}_2\text{CH}_2\text{COCH}_3}$$

When a ketone is to be alkylated, there may be two reactive carbon atoms available. It has been found that active methinyl groups are more readily alkylated than active methylene groups. An active methylene group bearing a phenyl group is more readily alkylated than one bearing only alkyl groups. The following examples illustrate these principles.25,31

²³ Boekelheide, J. Am. Chem. Soc., 69, 790 (1947).

³³³ Wilds and Shunk, J. Am. Chem. Soc., 72, 2388 (1950); see, however, Woodward et al., J. Am. Chem. Soc., 74, 4223 (1952). 205 Wilds and Werth, J. Org. Chem., 17, 1149, 1154 (1952).

n Crowley and Robinson, J. Chem. Soc., 1938, 2001.

In another useful version, the methiodide of 1-methyl-4-piperidone (XXXII), which may be considered as a Mannich base formed from two moles of formaldehyde and one mole each of acetone and methylamine, is used as an alkylating agent.³⁷ Only one of the carbon-nitrogen bonds breaks, and a 3-keto-5-dimethylaminoamyl group is thus introduced into the compound alkylated.

$$\begin{array}{c} O \\ & \downarrow \\ & \downarrow \\ N+I- \end{array} + CH_3COCH_2CO_2C_2H_5 \rightarrow \begin{array}{c} CH_3COCHCO_2C_2H_5 \\ & \downarrow \\ CH_2CH_2COCH_2CH_2N(CH_3)_2 \end{array}$$

The primary products may be capable of cyclization.37

Alkylation of Esters. Only esters containing doubly or triply activated carbon atoms have been alkylated by amine replacement reactions. Alkylations of α -nitro esters and β -keto esters have already been described.

Diethyl malonate has been monomethylated by means of tetramethylammonium ethoxide.38 Diethyl sodiomalonate has been benzylated, in yields as high as 79%, by means of quaternary salts containing, in addition to the benzyl group, methyl, ethyl, phenyl, or pentamethylene groups. Dibutyl ether, absolute ethanol, or an excess of diethyl malonate has been used as a solvent under various temperatures and pressures.7 Highest yields were obtained from diethyl sodiomalonate with benzyltrimethylammonium bromide in refluxing dibutyl ether (77%) or with benzyldimethylanilinium ehloride heated in the absence of solvent (73-79%). Diethyl sodiomalonate has also been alkylated with the methiodides of 1-dimethylaminomethyl-2-methoxynaphthalene 6 (IV, R = CH₃) and (+, -)-N,N-dimethyl- α -phenylethylamine,2 using Diethyl Carbitol as a solvent. When the methiodide of (+)-N,N-dimethyl-α-phenylethylamine (VI) was employed as an alkylating agent, the alkylation product was optically inactive; a small amount of N,N-dimethyl-α-phenylethylamine (probably formed by demethylation of the salt) was recovered from the reaction mixture and found to be only slightly optically active.2

Methyl eyanoaeetate and triearbethoxymethane have been benzylated with benzyldimethylamine.39 The initial step in this reaction is a

T Cardwell and McQuillin, J. Chem. Soc., 1949, 708.

Wittig, Heintzeler, and Wetterling, Ann., 557, 201 (1947).

²⁹ Snyder, Eliel, and Carnahan, J. Am. Chem. Soc., 72, 2958 (1950).

ethyl 36 or phenyl 30 groups, or phenols possessing meta bridges.27 Further examples of this type of alkylation are listed in Table VII.

$$\begin{array}{c} \text{CH}_3 \\ \text{CH}_2 = \text{C} \\ \text{CH}_3 \\ \text{$$

An interesting modification of this reaction consists in the use of the di-Mannich base of acctone; the simple alkylation product undergoes amine elimination to form a compound that can be cyclized to a dienonc capable of rearranging to a phenol.26,27,27a Whether an ortho- or metabridged phenol is obtained depends on the size of the alicyclic ring.276

$$(H_{3}C)_{3}N + H_{N}(CH_{3})_{3} + O = CO_{2}C_{2}H_{5}$$

$$CH_{2} + CH_{2} + O = CH_{2}$$

$$CH_{3} + CH_{2} + C$$

³⁵ Ghosh and Robinson, J. Chem. Soc., 1944, 506.

ethyl ³⁶ or phenyl ³⁰ groups, or phenols possessing meta bridges.²⁷ Further examples of this type of alkylation are listed in Table VII.

An interesting modification of this reaction consists in the use of the di-Mannich base of acetone; the simple alkylation product undergoes amine elimination to form a compound that can be cyclized to a dienone capable of rearranging to a phenol.^{26,27,27a} Whether an ortho- or metabridged phenol is obtained depends on the size of the alicyclic ring.^{27a}

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$$\begin{array}{c} O \\ \parallel \\ + CH_3COCH_2CO_2C_2H_5 \rightarrow \\ CH_3CH_2COCH_2CH_2CH_2N(CH_3)_2 \\ \\ CH_3 CH_3 \\ \times \times \times \times \times \times \\ \end{array}$$

The primary products may be capable of cyclization.37

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Methyl cyanoacetate and tricarbethoxymethane have been benzylated with benzyldimethylamine.39 The initial step in this reaction is a

Tardwell and McQuillin, J. Chem. Soc., 1949, 708.

³³ Wittig, Heintzeler, and Wetterling, Ann., 557, 201 (1947).

³³ Snyder, Eliel, and Carnahan, J. Am. Chem. Soc., 72, 2958 (1950).

(B)
$$\begin{array}{c} X \\ X \\ Y \\ Y \\ \end{array}$$

$$\begin{array}{c} X \\ CH_{2}N(CH_{3})_{2} + N_{a}CCO_{2}C_{2}H_{5} \xrightarrow{C_{2}H_{5}I} \\ Y \\ Y \\ \end{array}$$

$$\begin{array}{c} X \\ CH_{2}CCO_{2}C_{2}H_{5} + N_{a}I + \overset{+}{N}(CH_{3})_{2}(C_{2}H_{5})_{2}I - \overset{-}{N} \\ Y \\ \end{array}$$

$$\begin{array}{c} X \\ Y \\ \end{array}$$

The Mannich bases of indole, such as gramine (VIIa), have been used in alkylations of cyanoacetic and malonic esters. Yields of 85% were obtained by method A,⁷ whereas by method C a 76% yield was obtained in the alkylation of malonic ester.⁷ Tricarbethoxymethane, in the absence of added catalyst, has been alkylated by gramine (procedure C, 67% yield).¹⁷

1-Methylgramine (XXI) can be used as an alkylating agent for malonic ester derivatives (procedure C), although yields are low (9–15%); again the ester acts as a quaternizing agent in these reactions, since tertiary amines containing the alkyl group of the ester are formed.³⁹ Added base seems to decrease the rate of the reaction without appreciably reducing the yields. Higher yields are obtained by use of the methylgramine and the sodio derivative of the malonic methiodide of 1-methylgramine and the sodio derivative of the malonic ester; best yields are obtained with cyanomalonic ester (51%) and tricarbethoxymethane.¹⁷ In these last two reactions water may be used as a solvent since the active methylene compounds are more acidic than water.

(B)
$$\begin{array}{c} X \\ X \\ Y \\ Y \\ \\ X \\ X \\ X \\ X \\ X \\ \\ X \\$$

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or diethyl benzamidomalonate (XLIII).47 The methiodide of 1-methylgramine (XXI) reacts with the sodium salt of acetamidocyanoacetic ester (XLV); the product, obtained in 69% yield, can be hydrolyzed to 1-methyltryptophan (XLVI).48

Better yields of alkylation product are claimed when the quaternary salt is formed in situ (method B) by addition of two equivalents of ethyl iodide or dimethyl sulfate to a cooled mixture of the Mannich base with the sodio derivative of an amidomalonic ester in absolute ethanol.40 Thus, with gramine (VIIa) and ethyl acetamidocyanoacetate (XLV) or diethyl acetamidomalonate (XLI) yields of 98% and 95% have been reported. 49, 49, 50 Yields of 79-93% have been reported in alkylations of diethyl acetamidomalonate by this method with 2-, 4-, 5-, 6-, and 7-methylgramine.⁵¹ Ethyl acetamidocyanoacetate (XLV) was alkylated

⁴⁷ Albertson, Archer, and Suter, J. Am. Chem. Soc., 66, 500 (1944).

⁴⁸ Snyder and Eliel, J. Am. Chem. Soc., 70, 3855 (1948).

⁴⁹ Albertson and Tullar, J. Am. Chem. Soc., 67, 502 (1945). ⁵⁰ Albertson, Archer, and Suter, U. S. pats. 2,451,310 and 2,468,912 [C. A., 43, 1442,

⁵¹ Rydon, J. Chem. Soc., 1948, 705; Rydon and Siddapa, ibid., 1951, 2462; Kornfeld, 5806 (1949)]. J. Org. Chem., 16, 806 (1951); Hamlin and Fischer, J. Am. Chem. Soc., 73, 5007 (1951).

With 1-methylgramine (XXI), excess diethyl acetamidomalonate (XLI), and sodium, a total yield of only 12.5% of alkylation products is obtained.17

2-Dimethylaminomethylpyrrole reacts with diethyl acetamidomalonate (XLI) in toluene or xylene in the presence of sodium hydroxide to give a 70–80% yield of a product having the structure L^{43}

$$\begin{array}{c|c} CH_2N(C_2H_5)_2 & CH_2\\ NCO_2C_2H_6 & OCC_2C_2H_5\\ NHCOCH_3\\ L \end{array}$$

Diethyl malonate reacts slowly with Mannich bases of acetone 56 (XXIV) and cyclohexanone 57 at room temperature in ethanol containing small amounts of sodium ethoxide to form the normal simple alkylation products in yields of 43% and 86%, respectively. A rather low yield (16%) of ethyl 2-carbethoxy-5-ketohexanoate (LI) was obtained by reaction of diethyl sodiomalonate with the methiodide of 1-morpholino-3-butanone (LII).58 Powdered sodium hydroxide in xylene, as used in gramine alkylations (method C, p. 100), served as catalyst in an alkylation of diethyl malonate with a Mannich base of 2-phenylcyclohexanone (50% yield).58a

Reactions of ketonic Mannich bases with derivatives of aminomalonic ester or tricarbethoxymethane have not been reported. However, the following intramolecular reaction with a derivative of acetamidomalonic ester has led to a cyclopropane derivative. 585

$$(CH3)3NCH2CH2C(CO2C2H6)2 \rightarrow CH2 CO2C2H5 (33%)$$

$$OH^{-} NHCOCH3 \rightarrow CH2 - C-NHCOCH3$$

Methyl and ethyl cyanoacetate have been alkylated with the Mannich bases of 1-nitropropane, but the yields are not high (16-23%).21a

⁵⁶ Mannich and Fourneau, Ber., 71, 2090 (1938).

⁵⁸ Harradence and Lions, J. Proc. Roy. Soc. N. S. Wales, **72**, 233 (1939) [C. A., **33**, 5855

⁶⁵⁴ Bachmann and Wick, J. Am. Chem. Soc., 72, 3388 (1950).

⁵⁸⁵ Rinderknecht and Niemann, J. Am. Chem. Soc., 73, 4259 (1951).

by 3-diethylaminomethyl-5-methylindole (XLVII) in 87% yield by this method.⁵²

Yields of 90-94% were reported in alkylations by pyrrole Mannich bases of ethyl acetamidocyanoacetate (XLV) and diethyl acetamidomalonate (XLI), but a low yield was obtained with diethyl phthalimidomalonate (XLII).⁴³ Reaction of two moles of diethyl acetamidomalonate (XLI) with 2,5-bis(dimethylaminomethyl)pyrrole (XLVIII) occurs quantitatively by this method.⁴⁴

2-Acetamido-5-dimethylaminomethylthiazole (XXXVII, R = H) and the 4-methyl homolog (XXXVII, R = CH₃) have been used in alkylations of diethyl acetamidomalonate (XLI) in the presence of dimethyl sulfate. Of interest in this case is the use of the Mannich base hydrochloride, together with a molar excess of sodium ethoxide (to neutralize the hydrogen chloride).

Method C gives good yields in alkylations of aminomalonic ester derivatives with indole Mannich bases. Diethyl skatylacetamidomalonate (XLIV) is obtained in 90% yield when gramine (VIIa) and diethyl acetamidomalonate (XLI) are heated in xylene with powdered sodium hydroxide. Lower yields are obtained in pyridine, in the absence of a solvent, or in the absence of a eatalyst. Good to moderate yields are obtained when gramine (VIIa) is replaced by 3-diethylaminomethylindole (VII, $R = C_2H_5$) (85% yield) or 3-piperidinomethylindole (64%). Diethyl phthalimidomalonate (XLII) is alkylated to only a slight extent (10%) under the best of these conditions, but diethyl formamidomalonate gives the alkylation product in excellent yield (98%). Satisfactory yields of alkylation products have been obtained by this method in alkylations of diethyl acetamidomalonate (XLI) and ethyl acetamidocyanoacetate (XLV) with 5-bromogramine, 6-methylgramine. and 3-diethylaminomethyl-2-carbethoxyindole (XLIX).

With 1-methylgramine (XXI), excess diethyl acetamidomalonate (XLI), and sodium, a total yield of only 12.5% of alkylation products is obtained.¹⁷

2-Dimethylaminomethylpyrrole reacts with diethyl acetamidomalonate (XLI) in toluene or xylene in the presence of sodium hydroxide to give a 70-80% yield of a product having the structure L.⁴³

$$\begin{array}{c|c} CH_2N(C_2H_5)_2 & CH_2\\ CO_2C_2H_5 & CCO_2C_2H_5\\ H & NHCOCH_3 \end{array}$$

Diethyl malonate reacts slowly with Mannich bases of acetone ⁵⁶ (XXIV) and cyclohexanone ⁶⁷ at room temperature in ethanol containing small amounts of sodium ethoxide to form the normal simple alkylation products in yields of 43% and 86%, respectively. A rather low yield (16%) of ethyl 2-carbethoxy-5-ketohexanoate (LI) was obtained by reaction of diethyl sodiomalonate with the methiodide of 1-morpholino-3-butanone (LII). ⁵⁸ Powdered sodium hydroxide in xylene, as used in gramine alkylations (method C, p. 100), served as catalyst in an alkylation of diethyl malonate with a Mannich base of 2-phenylcyclohexanone (50% yield). ^{58a}

Reactions of ketonic Mannich bases with derivatives of aminomalonic ester or tricarbethoxymethane have not been reported. However, the following intramolecular reaction with a derivative of acetamidomalonic ester has led to a cyclopropane derivative.⁵⁸⁵

$$(CH_3)_3 \overset{+}{\text{NCH}}_2 CH_2 C(CO_2 C_2 H_5)_2 \longrightarrow CH_2 CO_2 C_2 H_5$$

$$OH^- \quad NHCOCH_3 \longrightarrow CH_2 -C-NHCOCH_3$$

$$(33\%)$$

Methyl and ethyl cyanoacetate have been alkylated with the Mannich bases of 1-nitropropane, but the yields are not high (16-23%).^{21a}

⁵⁶ Mannich and Fourneau, Ber., 71, 2090 (1938).

⁵⁷ Mannich and Koch. Ber., 75, 803 (1942).

⁵⁸ Harradence and Lions, J. Proc. Roy. Soc. N. S. Wales, 72, 233 (1939) [C. A., 33, 5855 (1939)].

bea Bachmann and Wick, J. Am. Chem. Soc., 72, 3388 (1950).

^{58b} Rinderknecht and Niemann, J. Am. Chem. Soc., 73, 4259 (1951).

An Alkylation of Indole

Indole reacts with diethyl piperidinomethylformamidomalonate to give diethyl skatylformamidomalonate 59 which is readily hydrolyzed to tryptophan in one step.52a The alkylation proceeds best in xylene

solution with a sodium hydroxide catalyst (76%); lower yields are obtained in other aromatic hydrocarbon solvents. In the absence of the basic catalyst, 3-piperidinomethylindole (VII, R_2 = pentamethylene) is the principal or exclusive product. Other alkylations with Mannich bases of formamidomalonic ester have been reported. 59a, b Indole has also been alkylated with diethylaminoacetonitrile.59c

Amine Replacement Reactions of Quaternary Salts with Organometallic Compounds

Only a few reactions of Grignard and organolithium reagents with quaternary ammonium salts resulting in displacement of the ammonium nitrogen by the alkyl group of the organometallic reagent are on record. The reaction apparently has not been studied extensively. 9-Fluoryllithium reacts with tetramethylammonium chloride to yield 9-methylfluorene in unspecified yield.38 Phenyllithium reacts in a different fashion.60 From the reaction of phenyllithium with benzyltrimethylammonium bromide, no diphenylmethane was isolated; the latter was apparently metallated as formed and further alkylated by the quaternary salt to 1,1,2-triphenylethane. α-Phenylethyldimethylamine was

⁵⁹ Butenandt, Hellmann, and Renz, Z. physiol. Chem., 284, 175 (1949); C. Y. Meyers, doctoral thesis, University of Illinois, Urbana, Ill., 1951.

^{69a} Hellmann and Brendle, Z. physiol. Chem., 287, 235 (1951).

⁶⁹b Hellmann and Renz, Chem. Ber., 84, 901 (1951).

^{69c} N. J. Murphy, bachelor's thesis, University of Notre Dame, Notre Dame, Ind., 1952. 60 Wittig and co-workers, Ann., 555, 133 (1944); 557, 193 (1947). For a review see: Wittig, Angew. Chem., 63, 15 (1951).

also obtained.⁶¹ The methiodide of 1-methylgramine (XXI) reacts with methylmagnesium iodide and with phenylmagnesium bromide in refluxing dibutyl ether to yield 1-methyl-3-ethylindole (LIII) and

$$\begin{array}{c} C_{6}H_{5}Li + C_{6}H_{5}CH_{2}\overset{+}{N}(CH_{3})_{3} \; Br^{-} \rightarrow LiBr + N(CH_{3})_{3} + C_{6}H_{5}CH_{2}C_{6}H_{5} \\ \\ C_{6}H_{5}CH_{2}C_{6}H_{5} + C_{6}H_{6}Li \rightarrow C_{6}H_{5} + C_{6}H_{5}CHLiC_{6}H_{5} \end{array}$$

$$C_6H_6CHLiC_6H_5 + C_6H_6CH_2\overset{+}{N}(CH_3)_3 Br^- \rightarrow$$

$$LiBr + N(CH_3)_3 + C_6H_5CH_2CH(C_6H_6)_2$$

1-methyl-3-benzylindole (LIV).⁶² The methiodide of gramine (VIIa) similarly yields 3-ethylindole (LV), 3-benzylindole (LVI), and 3-phenethylindole (LVII), although in poor yield; a by-product with the composition and properties of sym-3,3-diindolylethane (LVIII) is presumably formed by a coupling reaction (equation on p. 133). 3-Benzylindole was obtained in only 3% yield when the tertiary amine gramine was treated with phenylmagnesium bromide. Attempts to extend the reaction with organometallic reagents to a number of other Mannich bases and quaternary salts were unsuccessful.⁶² N,N'-Benzaldipiperidine (LIX,

$$\mathbb{R}_{1}^{\mathbb{R}_{2}}$$

LIII $R_1 = CH_3$, $R_2 = C_2H_5$ LIV $R_1 = CH_3$, $R_2 = C_6H_5CH_2$ LV $R_1 = H$, $R_2 = C_2H_5$ LVI $R_1 = H$, $R_2 = C_6H_5CH_2$ LVII $R_1 = H$, $R_2 = C_6H_5CH_2CH_2$

R=H) and N,N'-benzaldi- γ -pipecoline (LIX, $R=CH_3$) react with benzylmagnesium chloride to give 1-piperidino-1,2-diphenylethane (LX, R=H) and 1-(γ -pipecolino)-1,2-diphenylethane (LX, $R=CH_3$)

in 18 and 14% yield, respectively.63

⁶¹ Wittig, Mangold, and Felletschin, Ann., 560, 116 (1948).

Snyder, Eliel, and Carnahan, J. Am. Chem. Soc., 73, 970 (1951).
 Goodson and Christopher, J. Am. Chem. Soc., 72, 358 (1950).

$$C_6H_5CH$$
 $+ C_6H_5CH_2MgCl \rightarrow$
 R
 $+ C_6H_5CH_2MgCl \rightarrow$
 R
 C_6H_6CH
 $+ R$
 $NMgCl$
 $CH_2C_6H_5$

MECHANISM OF THE REACTION

The path by which alkylations with tertiary amines and quaternary ammonium salts proceed has not yet been definitely established, and any statements concerning the mechanism of the reaction are therefore speculative.

Alkylations with Tertiary Amines

The mechanism that has most frequently been proposed for alkylations with tertiary amines involves the elimination of a secondary amine, resulting in the formation of an unsaturated compound which undergoes addition of the species to be alkylated.

$$ACH_2CH_2NR_2 \rightarrow NHR_2 + ACH \rightleftharpoons CH_2$$

 $ACH \rightleftharpoons CH_2 + CHRR'R'' \rightarrow ACH_2CH_2CRR'R''$

A scheme of this type was first proposed for alkylations with phenolic Mannich bases by von Auwers.⁶⁴⁻⁶⁸ The hypothetical intermediate is a methylenequinone whose formation involves 1,4- or 1,6-elimination.

$$\begin{array}{c}
\text{OH} \\
\text{CH}_2\text{NR}_2 \\
\text{OH}
\end{array}$$

$$\begin{array}{c}
\text{OH} \\
\text{CH}_2\text{CRR'R''}
\end{array}$$

68 Dalgliesh, J. Am. Chem. Soc., 71, 1697 (1949).

⁶⁴ v. Auwers, Ber., 36, 1878 (1903).

⁶⁵ v. Auwers, Ann., 344, 131 (1906).

⁶⁶ v. Auwers and Bullmann, Ber., 59, 2719 (1926).

⁶⁷ Snyder and Brewster, J. Am. Chem. Soc., 70, 4230 (1948).

$$\begin{array}{c}
\text{OH} \\
\text{OH} \\
\text{CH}_2\text{NR}_2
\end{array}
\rightarrow \text{HNR}_2 +
\begin{array}{c}
\text{O} \\
\text{CHRR'R''}
\end{array}$$

$$\begin{array}{c}
\text{CH}_2\text{CRR'R''}
\end{array}$$

A similar scheme has been proposed 9 for alkylations with gramine (VIIa).

$$\begin{array}{c} CH_2N(CH_3)_2 \\ \\ N \\ \\ VIIa \\ \\ HN(CH_3)_2 + \\ \\ N \end{array} \longrightarrow \begin{array}{c} CH_2 \\ \\ \\ N \end{array} \longrightarrow \begin{array}{c} CH_2CN \\ \\ \\ N \\ \\ H \end{array}$$

1,2-Elimination may be the first step in alkylations with ketonic Mannich bases.24

For the ketonic Mannich bases, the elimination-addition mechanism is supported by the facts that these compounds will yield α,β -unsaturated ketones by elimination of secondary amines $^{61, \, 69, \, 70, \, 71}$ and that α, β unsaturated ketones will add active methylene compounds (Michael reaction).

The elimination of the secondary amine may be either an acidcatalyzed E_1 (mechanism A) or a base-catalyzed E_2 (mechanism B) reaction.72 In the simple elimination reactions of ketonic Mannich

⁶⁹ Mannich and co-workers, Ber., 53, 1374 (1920); 55, 356, 3510 (1922); 57, 1116 (1924);

⁷⁰ Mannich and Hönig, Arch. Pharm., 265, 598 (1927). ⁷¹ Harradence and Lions, J. Proc. Roy. Soc. N. S. Wales, 72, 284 (1939) [C. A., 33, 6825]

Remick, Electronic Interpretations of Organic Chemistry, 2nd ed., p. 424, John Wiley & Sons, 1949.

(A)
$$ACH_2CH_2NR_2 + H^+ \rightarrow ACH_2CH_2NR_2H^+ \rightarrow NHR_2 + ACH_2CH_2^+ \rightarrow ACH = CH_2 + H^+$$

(B)
$$ACH_2CH_2NR_2 + B$$
: $\rightarrow A\overline{C}HCH_2NR_2 + BH^+$
 $A\overline{C}HCH_2NR_2 \rightarrow ACH = CH_2 + NR_2^-$
 $NR_2^- + BH^+ \rightarrow NHR_2 + B$:

bases, both acid catalysis 69,70 and base catalysis 61,73 have been observed. It is also possible that reaction occurs between two molecules of the Mannich base, one acting as an acid and the other as a base. Still another possibility with ketonic and ortho-substituted phenolic Mannich bases is an intramolecular elimination involving a chelate intermediate.

Only the enolic form of a ketonic Mannich base is capable of chelation.

$$\begin{array}{c} \text{CH} \\ \text{RC} \\ \downarrow \\ \text{O} \\ \text{NR}_2 \end{array} \rightarrow \begin{array}{c} \text{R-C=CH-CH}_2^+ \leftrightarrow \text{R-C-CH=CH}_2 + \text{NHR}_2 \\ \downarrow \\ \text{O-} \end{array}$$

An attempt to obtain spectral evidence for the existence of this type of intermediate has, however, failed.74

The Michael addition of an active methylene compound to an activated unsaturated species is known to be base catalyzed. The over-all alkylation reaction would therefore be expected to be either base or acid-base catalyzed, and this is actually found to be so. Since one of the reactants is itself quite basic, the addition of an extrinsic basic catalyst is sometimes unnecessary or even undesirable.^{17, 19,20} In the alkylation of dibenzoylmethane by 1-morpholinomethyl-2-naphthol (XXVII), the reaction is known to be catalyzed by added hydrochloric acid.²³

The facts that benzyldimethylamine and 1-methylgramine (XXI) will alkylate methyl cyanoacetate and tricarbethoxymethane and that 1-methylgramine will alkylate diethyl acetamidomalonate (XLI),^{17,39} although these amines are structurally incapable of reacting by an

⁷³ Bruylants, Bull. soc. chim. Belg., 32, 256 (1923).

⁷⁴ Brewster, unpublished observations.

elimination-addition mechanism, have been satisfactorily explained by demonstrating that alkylation is preceded by quaternization ³⁹ (p. 118). However, 1-methylgramine (XXI) also alkylates secondary amines 75 and 1-methylindole,17 and these reactions (like the reaction of 2-dimethylaminomethyl-2-nitropropane with piperidine 21a) cannot be explained as alkylations with quaternary salts; they will take place only in the presence of acids 75 and might therefore proceed by a path resembling that of mechanism A above (p. 128). It should be noted that one of the intermediates in this mechanism is a carbonium ion and that the loss of a proton from this ion to form the unsaturated compound is not essential, since the carbonium ion itself could be the alkylating agent.

essential, since the Carbonian
$$CH_2N(CH_3)_2$$
 + $H^+ \rightarrow CH_3$ $CH_2NH(CH_3)_2^+ \rightarrow CH_3$ CH_3 CH_3 CH_3 $CH_2^+ + CRR'R'' \rightarrow CH_3$ CH_3

One would expect the carbonium ion postulated in this mechanism to be stabilized by resonance.

Other types of Mannich bases may react by the same path.

Another possible path for the alkylation reactions with 1-methylgramine hydrochloride, the hydrochloride of 2-dimethylaminomethyl-2-nitropropane (p. 139) and the Mannich base of diethyl formamidomalonate (p. 124), none of which can react by elimination-addition, is a complete reversal of the Mannich reaction, 58a, 68, 76, 77 followed by re-

⁷⁵ Snyder and Eliel, J. Am. Chem. Soc., 70, 4233 (1948).

⁷⁶ Mannich and Kather, Arch. Pharm., 257, 18 (1919).

⁷ Kermack and Muir, J. Chem. Soc., 1931, 3089.

combination of the fragments. This may also be the path of alkylations with diethylaminoacetonitrile. 59a,c

$$\begin{array}{c} CH_2N(CH_3)_2 + H_2O \xrightarrow{H^+} NH(CH_3)_2 + CH_2O + \\ N \\ CH_3 \end{array}$$

$$\begin{array}{c} CH_3 \\ CH_3 \end{array} + CH_2O \xrightarrow{H^+} CH_3 \xrightarrow{CH_3} CH_3$$

$$\begin{array}{c} CH_2NC_5H_{10} \\ CH_3 \end{array}$$

$$\begin{array}{c} CH_2NC_5H_{10} \\ CH_3 \end{array}$$

As a further possibility, alkylation reactions with tertiary amines may involve a nucleophilic displacement. Such a path seems less likely in

$$\begin{array}{c} \operatorname{RCH_2N}(\operatorname{CH_3})_2{}^+ + \operatorname{B}{}^- \to \operatorname{RCH_2B} + \operatorname{NH}(\operatorname{CH_3})_2 \\ \text{H} \end{array}$$

view of the fact that the base would be expected to abstract a proton from the ammonium salt rather than displace a dimethylamine molecule.

Alkylations with Quaternary Salts

It has been proposed that alkylations with quaternary salts of ketonic Mannich bases proceed by the same elimination-addition mechanism as alkylations with the Mannich bases themselves. The elimination step might be of the E_1 type (loss of a tertiary amine followed by loss of a proton) or of the E_2 type (abstraction of a proton followed by loss of a tertiary amine). β -Dimethylaminopivalophenone (XV), a ketonic Mannich base that is structurally incapable of undergoing amine elimination, will not act as an alkylating agent. On the other hand there are numerous quaternary ammonium salts that act as alkylating agents although they show no tendency to undergo amine elimination, viz., quaternary salts of benzyldialkylamines, 4.6.7.39 substituted benzyldialkylamines, 2.6 and 1-methylgramine (XXI).9.17.48 It therefore appears that elimination-addition is not the only path by which alkylation reactions

with quaternary bases may proceed, the alternative being direct substitution. It might be noted that β -dimethylaminopival ophenone (XV) is an amine of the neopentyl type and would therefore not be expected to undergo bimolecular substitution reactions readily.

The question whether the substitution is of the S_N1 or S_N2 type 78 has not been answered definitely for carbon-carbon alkylations. It has been found that the pyrolysis of (+)-α-phenylethyltrimethylammonium acetate to α -phenylethyl acetate proceeds with complete or almost complete inversion,2 but in carbon alkylations with the active quaternary iodide, VI, both the product and recovered starting material were racemized.2 Thus, although the reaction of the quaternary acetate is of the S_N2 type, no conclusions can be arrived at with regard to the mechanism of the carbon alkylation since racemization may have been due to abstruction of a proton from the α -carbon by the basic catalyst with concomitant loss of asymmetry. Dipolar ions of the type represented by LXI and known as "alkylides" have been observed in other in-

$$C_6H_6CHN(CH_3)_3^+ + B: \rightarrow C_6H_6CN(CH_3)_3^+ + B:H$$
 CH_3
 CH_3
 LXI

stances; 38,60 a similar ion is probably responsible for the racemization of optically active nicotine dimethiodide (LXII) by aqueous base at 100°,61,79

Allylic rearrangements have been observed in alkylations of sodium cyanide with the methiodide of 1-methylgramine (IX) 9 (p. 107) and furfuryltrimethylammonium iodide 10 (p. 107). It is of interest that the ratio of rearranged to normal product in the latter reaction is much smaller than in the alkylation of sodium cyanide with furfuryl chloride. 80, 81 Whereas it formerly was thought that allylic rearrangements were indicative of carbonium-ion intermediates, it is now recognized that they may occur even in reactions that are subject to second-order

⁷⁸ See ref. 72, p. 74.

⁷⁹ Späth and Bobenberger, Ber., 77, 362 (1944).

⁸⁰ Runde, Scott, and Johnson, J. Am. Chem. Soc., 52, 1284 (1930).

⁸¹ Reichstein, Ber., 63, 749 (1930).

kinetics.⁸² Therefore the occurrence of such rearrangements in alkylations with quaternary ammonium salts is not necessarily indicative of an $S_N 1$ (carbonium ion) mechanism.

Further experimentation is needed for definite elucidation of the exact mechanism by which these reactions proceed.

RELATED REACTIONS

It seems desirable, for the sake of completeness, to describe briefly the more important reactions of carbon, nitrogen, oxygen, sulfur, and halogen alkylation by amine replacement, which for various reasons have not been considered in detail in the preceding sections and are omitted from the tables. The following résumé does not pretend to be complete, and only leading references are listed.

Carbon-Carbon Alkylations

The carbon-carbon alkylation reactions of labile amino compounds that were not reviewed in detail fall into the following five categories: (a) those in which intermolecular "self-alkylation" occurs; (b) those in which intramolecular "self-alkylation" or rearrangement occurs; (c) those in which the carbon-nitrogen bond broken is one of the bonds of a heteroaromatic system; (d) those in which the carbon-nitrogen bond broken is found in a diaminomethane; (e) those in which the new carbon-carbon bond formed is part of an ethylenic double bond. Examples of each of the more important types of these reactions are given below.

Intermolecular Self-Alkylations. Self-Alkylation of Phenolic and Indole Mannich Bases. Auwers and his co-workers ^{14, 64, 66, 83-85} found that o- and p-hydroxybenzylamines (many of which cannot be made by the Mannich reaction) readily form diarylmethanes by the loss of formal-dehyde and two moles of amine in weakly alkaline solution, according to the equation on p. 109. This reaction is prominent in attempts to use phenolic Mannich bases as alkylating agents. ^{12, 16} A similar reaction occurs when 1-methylgramine (XXI) is used in alkylations of malonic ester derivatives or when the hydrochloride or methiodide of 1-methylgramine is heated in dilute aqueous alkali. ¹⁷ The Mannich bases ob-

E Kepner, Winstein, and Young. J. Am. Chem. Soc., 71, 115 (1949).

⁸³ v. Auwers and Senter, Ber., 29, 1120 (1896).

v. Auwers and co-workers, Ber., 28, 2910 (1895); 29, 1110 (1896).
 v. Auwers and co-workers, Ann., 344, 141, 171, 194, 227, 257 (1906).

tained by condensing indoles, benzaldehyde, and aromatic amines undergo similar reactions when heated with dilute hydrochloric acid. 86, 87, 88

Self-Alkylation of 9-Fluoryltrimethylammonium Hydroxide. Trimethylfluorylammonium hydroxide forms, among other products, dibiphenyleneethylene when heated.89 The hydrogen atom at the 9 position of the fluorene residue is activated by two aromatic residues and a quaternary ammonium grouping; this hydrogen atom is probably replaced in an alkylation process. The primary product formed by such a reaction is a quaternary ammonium hydroxide, which would be expected to undergo a particularly easy amine elimination. (See ref. 91a for a similar reaction.)

Coupling of Quaternary Ammonium Salts. When quaternary salts of gramine 62 (VIIa) or benzhydryldimethylamine 61 are treated with organometallic reagents, one of the reactions that occurs is coupling of the reactive alkyl residues of the amines.

$$2RN(CH_3)_3 X^- + 2R'M \rightarrow R-R+R'-R'+2N(CH_3)_3 + 2MX$$
 $R = benzhydryl or skatyl$

This reaction resembles the coupling of benzyl halides by Grignard reagents.

⁸⁶ Passerini and Bonciani, Gazz. chim. ital., 63, 138 (1933).

⁸⁷ Passerini and Albani, Gazz. chim. ital., 65, 933 (1935).

⁸³ Neri, Gazz. chim. ital., 64, 420 (1934).

⁸⁹ Ingold and Jessop, J. Chem. Soc., 1929, 2357; 1930, 713.

Reductive Coupling of Ethanolamines. This rather specific reaction was discovered by Wittig and co-workers. 61

$$2(C_6H_5)_2COHCH_2N(CH_3)_2 + 6K \rightarrow$$

$$(C_6H_6)_2CHCH_2CH_2CH(C_6H_6)_2 + 2K_2O + 2KN(CH_3)_2$$

Intramolecular Self-Alkylations. The Stevens Rearrangement. 60, 61, 89a-91

The Sommelet Rearrangement. 61, 91a, 92

The Hofmann-Martius Rearrangement. 93, 94, 95

Some quaternary salts of phenolic Mannich bases, in which the amino group is present in an aniline derivative, rearrange readily in alkaline solution to form substituted benzylanilines. 56, 83, 96, 97

^{89a} Stevens and co-workers, J. Chem. Soc., 1928, 3193; 1930, 2107, 2119; 1932, 55, 1926, 1932; 1934, 279.

⁹⁰ Campbell, Houston, and Kenyon, J. Chem. Soc., 1947, 93. Bock, Smith, and Auten, Atlantic City Meeting of the American Chemical Society, 1949, Abstracts, p. 70M.

⁹¹ Dahn and Solms, Helv. Chim. Acla, 34, 907 (1951); Brewster and Kline, J. Am. Chem. Soc., 74, 5179 (1952).

¹ Kantor and Hauser, J. Am. Chem. Soc., 73, 4122 (1951).

⁵² Sommelet, Compt. rend., 205, 56 (1937).

⁹³ Hickinbottom and Ryder, J. Chem. Soc., 1931, 1281.

⁴ Hey, J. Chem. Soc., 1931, 1581.

²⁵ Wittig and Merkle, Ber., 76, 109 (1943).

Zincke and Hunke, Ann., 349, 83 (1906); v. Auwers, Ann., 334, 264 (1904).
 Corley and Blout, J. Am. Chem. Soc., 59, 761 (1947).

The Ladenburg Rearrangement. 98, 99

The Rearrangement of Diacylanilines. 100

$$N(COCH_3)_2 \xrightarrow{Z_nCl_2} CH_3CO$$
NHCOCH

Reactions in Which the Carbon-Nitrogen Bond Broken Is One of the Bonds of a Heteroaromatic System. The Reissert Reaction, 101, 102

$$+ \text{RCOCl} + \text{KCN} \rightarrow \text{CN} + \text{KCl}$$

The products of this reaction (so-called Reissert compounds) are usually employed in the synthesis of aldehydes.

$$\begin{array}{c} H \\ \downarrow \\ \text{COR} \end{array} + 2H_2O \rightarrow \begin{array}{c} \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \end{array}$$

⁹⁸ Ladenburg, Ber., 16, 1410, 2057 (1883); Ann., 247, 1 (1888).

⁹⁹ Crook, J. Am. Chem. Soc., 70, 416 (1948).

¹⁰⁰ Chapman, J. Chem. Soc., 127, 2818 (1925). Chapman, J. Chem. Soc., 121, 2010 (1995); Sugasawa and Tsuda, J. Pharm. Soc., Japan, 56, 103 (1905); Reissert, Ber., 38, 1603 (1905); Carabainta and Wischer, J. Am. Ch. Reissert, Ber., 38, 1603 (1903), Sugaranta and Fischer, J. Am. Chem. Soc., 63, 2021 (1936) [C. A., 32, 5836 (1938)]; Grosheintz and Fischer, J. Am. Chem. Soc., 63, 2021 (1936) [C. A., 32, 5836 (1938)]; Grosnentz Land Hazlett, ibid., 71, 1949 (1949). (1941); Woodward, ibid., 62, 1626 (1940); McEwen and Hazlett, ibid., 71, 1949 (1949). 102 Manske, Chem. Revs., 30, 113, 145 (1942).

The Reissert compounds may also be alkylated by Mannich bases. 102a

The Reaction of Alkali Cyanides with Alkylpyridinium Salts. 102, 103

The Reaction of Nitro Compounds with Alkylpyridinium Salts. 104

103 Kaufmann, Ber., 51, 116 (1918); Leonard and Foster, J. Am. Chem. Soc., 74, 2110, 3671 (1952).

Leonard and Leubner, J. Am. Chem. Soc., 71, 3405 (1949); Leonard, Leubner, and Burk, J. Org. Chem., 15, 979 (1950); Leonard, DeWalt, and Leubner, J. Am. Chem. Soc.,

Nitrogen Alkylations

Amine Exchange Reactions of Quaternary Salts. When many quaternary ammonium salts, particularly those containing benzyl, allyl, or methyl groups, are heated with ammonia or with primary or secondary amines, an exchange of amino groups takes place. 10, 15, 16, 111, 112, 113

$$\begin{bmatrix} R \\ | \\ R'-N-R \\ | \\ R \end{bmatrix}^+ + HNR''_2 \rightarrow R'NR''_2 + NR_3 + H^+$$

Amine Exchange Reactions of Mannich Bases. Simple amine exchange reactions have been observed with Mannich bases of nitroalkanes, 21a,114 indole 41 (VII), phenols, 15a and ketones, 67,115 as well as with the benzaldehyde Mannich bases of β -naphthol 67 (LXIII).

Quaternary salts of some Mannich bases (e.g., those of indole, VII, and those of acetophenone, XXV) react readily by amine exchange with tertiary amines (including Mannich bases) to give new quaternary salts. This reaction may be important as a side reaction in the quaternization of Mannich bases by means of such reagents as methyl iodide, ⁶⁵ for example, in the quaternization of gramine.

$$3 \underbrace{\begin{array}{c} CH_2N(CH_3)_2 \\ H \end{array}}_{V11\alpha} + 3CH_3I \rightarrow \underbrace{\begin{array}{c} CH_2N(CH_3)_3 \\ H \end{array}}_{N}$$

¹¹¹ Scholtz, Ber., 24, 2402 (1891); 31, 414, 1700 (1898).

¹¹² v. Braun and co-workers, Ann., 445, 247 (1925); Ber., 59, 1786, 2330 (1926).

¹¹³ Hultquist and co-workers, J. Am. Chem. Soc., 70, 23 (1948).
114 Duden, Bock, and Reid, Ber., 38, 2036 (1905).

¹¹⁶ Denton, Schedl, Neier, and Brookfield, J. Am. Chem. Soc., 72, 3792 (1950).

Compounds that do not permit amine elimination, such as α -dimethylaminomethyl- β -methoxynaphthalene ² (IV, R = CH₃), β -dimethylaminopivalophenone 11 (XV), 1-methylgramine 75 (XXI), and 2-dimethylaminomethyl-2-nitropropane 21a do not undergo an amine exchange reaction in the absence of added acid catalyst, such as hydrogen chloride or boron trifluoride.75

Formation of Pyrazolines from Ketonic Mannich Bases. The phenylhydrazones of ketonic Mannich bases form pyrazolines by internal amine exchange under conditions similar to those required for phenylhydrazone formation. 1, 116-120

Conversion of Mannich Bases into Aldehydes. In an extension of the amine exchange reactions of Mannich bases, the base in acetic acid solution is allowed to react with hexamethylene tetramine. 121 intermediate quaternary salt decomposes to yield an aldehyde.

$$RCH_2N(CH_3)_2 + (CH_2)_6N_4 + CH_3CO_2H \rightarrow$$

$$NH(CH_3)_2 + RCH_2N(CH_2)_6N_3^+ + CH_3CO_2^- \rightarrow RCHO$$

This process, which resembles the Sommelet reaction 122 for converting benzyl halides into aromatic aldehydes, has been applied successfully to the Mannich bases of indole, 2-phenylindole, 2-carbethoxyindole, phenol, and β -naphthol, but has failed with Mannich bases of acetophenone, pyrrole, and 2-nitro-3-methylthiophene as well as 2-nitropropane. It was successful also with benzylamine and N-methylbenzylamine, but not with N,N-dimethylbenzylamine.

¹¹⁶ Mannieh and Bauroth, Ber., 57, 1108 (1924).

¹¹⁷ Nisbet and Gray, J. Chem. Soc., 1933, 839.

¹¹⁸ Levvy and Nisbet, J. Chem. Soc., 1938, 1053, 1572.

¹¹⁹ Nisbet, J. Chem. Soc., 1938, 1237, 1568; 1945, 126.

¹²⁰ Harradenee and Lions, J. Proc. Roy. Soc. N. S. Wales, 73, 14 (1939) [C. A., 33, 8196 (1939)].

¹²¹ Snyder, Swaminathan, and Sims, J. Am. Chem. Soc., 74, 5110 (1952).

¹²² Sommelet, Compt. rend., 157, 852 (1913); Angyal and co-workers, J. Chem. Soc., 1949, 2700, 2704; 1950, 2141.

Oxygen Alkylations

Formation of Alcohols from Quaternary Ammonium Hydroxides. Quaternary ammonium hydroxides, when heated strongly, may form alcohols rather than olefins, particularly when benzyl, allyl, or, in some cases, methyl groups are present and when no radicals, such as ethyl or phenethyl, that lead to easy formation of olefins are present. 123-127

$$R'NR_3OH^- \rightarrow R'OH + NR_3$$

The formation of pseudobases from pyridinium hydroxides is formally similar to the formation of alcohols from quaternary ammonium hydroxides.^{128,129}

Formation of Ethers from Quaternary Ammonium Phenoxides. 4, 130-136a Quaternary ammonium compounds have been used in the formation of benzyl, methyl, ethyl, and allyl ethers of phenols.

Some of the quaternary ethoxides of p-nitroaniline and p-formylaniline (p-aminobenzaldehyde) decompose to form alkoxy substituted benzenes.¹³⁷

Epoxides are formed in the Hofmann degradation of quaternary salts of 1-hydroxy-2-amines. 138, 139, 140

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    Hofmann, Ann., 78, 253 (1851); 79, 11 (1851); Ber., 14, 494 (1881).
    Ingold and Vass, J. Chem. Soc., 1928, 3125.
    von Braun, Teuffert, and Weissbach, Ann., 472, 121 (1929).
    Hanhart and Ingold, J. Chem. Soc., 1927, 997.
    von Braun, Ann., 382, 1 (1911).
    Decker, Ber., 25, 443 (1892).
    Hantzsch and Kalb, Ber., 32, 3109 (1899).
    Haneley and Turner, J. Chem. Soc., 3, 101 (1926) [C. A., 20, 3695 (1926)].
    Henley and Turner, J. Chem. Soc., 1931, 1172.
    Griess, Ber., 13, 246 (1880).
    Boehringer, Ger. pat. 247,180 [Frdl., 10, 1215 (1912)].
    Rodionow, Bull. soc. chim. France, [4] 39, 305 (1926).
    Rodionow, Bull. soc. chim. France, [4] 45, 109 (1929).
    Tarbell and Vaughan, J. Am. Chem. Soc., 65, 231 (1943).
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135a Kursanow, Setkina, and Rodionow, Bull. acad. sci. U.R.S.S., Classe sci. chim., 1948, 228 [C. A., 42, 4922 (1948)]; Kursanow and Setkina, Doklady Akad. Nauk S.S.S.R., 65, 847 (1949) [C. A., 43, 6622c (1949)]; Setkina and Kursanow, Izvest. Akad. Nauk S.S.S.R., Otdel. Khim. Nauk, 1949, 311 [C. A., 44, 159a (1950)]; ibid., 1951, 81 [C. A., 46, 458 (1952)]; Setkina, Izvest. Akad. Nauk S.S.S.R., Otdel. Khim. Nauk, 1950, 216 [C. A., 44, 9337c (1950)].

Zaki and Fahim, J. Chem. Soc., 1942, 270; Zaki and Tadros, J. Chem. Soc., 1941, 350.
 von Braun and Schirmacher. Ber., 55, 1845 (1923).

123 von Braun, Ber., 56, 2178 (1923).

140 von Braun and Münch, Ber., 59, 1941 (1926); Curtin, Harris, and Pollak, J. Am. Chem. Soc., 73, 3453 (1951).

Formation of Esters from Quaternary Ammonium Salts of Carboxylic Acids. Benzyl 4 and methyl 141 and ethyl 141a esters of carboxylic acids have been prepared by heating the acids with quaternary ammonium hydroxides containing the appropriate radicals as the most readily replaced substituents on the nitrogen atom. Benzyl esters may also be obtained by heating methyl esters with benzyldimethylamine.142

Benzyldimethylamine reacts with acetic anhydride or benzoyl chloride to give benzyl acetate and benzoate respectively.143 Phenolic Mannich bases similarly form acetyl derivatives of the corresponding methylolphenols.^{14, 15a, 144, 145, 145a}

OH
$$CH_2N(CH_3)_2 + 2(CH_3CO)_2O \rightarrow$$

$$OCOCH_3$$

$$CH_2OCOCH_3 + CH_3CON(CH_3)_2 + CH_3CO_2H$$
Sulfur Alkylations

Quaternary ammonium salts containing such anions as sulfide, hydrosulfide, mercaptide, 5, 146, 147 thiosulfate, thiocyanate, bisulfite, sulfite, 5,147 and p-toluenesulfinate 4 decompose when heated to form alkyl derivatives of these anions containing carbon-sulfur bonds. Alkyl groups that can take part easily in these reactions are allyl,147 benzyl,5 and methyl,146 in order of decreasing activity.

The ready cleavage of thiamin by bisulfite ion indicated the presence of a reactive benzyl type of quaternary ammonium group in the molecule. 148

Among tertiary amines, gramine (VIIa) has been used in the alkylation of sodium bisulfite. 148a The Mannich bases of phenol will alkylate mercaptans. 1486 An extensive study of sulfur alkylations has been reported.21b

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141 Lawson and Collie, J. Chem. Soc., 53, 624 (1888); Prelog and Piantanida, Z. physiol.
Chem., 244, 56 (1936); Fuson, Corse, and Horning, J. Am. Chem. Soc., 61, 1290 (1939).
  141a Kupferberg, J. prakt. Chem., [2] 16, 440 (1877).
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¹¹² Eliel and Anderson, J. Am. Chem. Soc., 74, 547 (1952).

¹⁰ Tiffeneau and Fuhrer, Bull. soc. chim. France, (4) 15, 162 (1914). 144 Madinaveitia, Anales soc. españ. fís. y quím., 19, 259 (1921) [C. A., 16, 1230 (1922)].

¹¹⁵ Bruson and MacMullen, J. Am. Chem. Soc., 63, 270 (1941). 1130 For similar reactions, see Setkina and Kursanow, Izrest. Akad. Nauk S.S.S.R., Otdel. Khim. Nauk, 1949, 190 [C. A., 43, 6161h (1949)].

¹⁴⁶ Clarke, J. Chem. Soc., 103, 1689 (1913).

¹⁰ Snyder and Speek, J. Am. Chem. Soc., 61, 2895 (1939). 16 Williams, Waterman, Keresztesy, and Buchman, J. Am. Chem. Soc., 57, 536 (1935).

¹⁸³⁴ Wieland, Fischer, and Moewus, Ann., 551, 47 (1948). 165 McCleary and Roberts, U. S. pat. 2,417,118 [C. A., 41, 3819b (1947)].

Halogen Alkylations

Decomposition of Quaternary Ammonium Halides. ammonium halides decompose when heated to form alkyl halides and tertiary amines. 123 Mixtures of amines and halides are often obtained

$$\begin{bmatrix} R \\ | \\ R'-N-R \\ | \\ R \end{bmatrix}^+ X^- \to R'X + NR_3$$

from mixed quaternary halides. Allyl, benzyl, and methyl start and methyl start. groups are lost as halides more readily than are other alkyl groups or the phenyl group.¹²⁷ Quaternary ammonium halides containing an asymmetric nitrogen atom racemize readily in solution at room temperature. 154

The von Braun Cyanogen Bromide Reaction. Cyanogen bromide reacts with a tertiary amine to form a quaternary salt, which readily decomposes to form an alkyl halide and a dialkylcyanamide. 155, 156

$$\begin{array}{c} R \\ | \\ R'N + BrCN \\ | \\ R \end{array} \rightarrow \begin{bmatrix} R \\ | \\ R'NCN \\ | \\ R \end{bmatrix}^+ Br^- \rightarrow R'Br + NCN \\ | \\ R \end{array}$$

This reaction was extensively studied by von Braun.¹⁶⁷ Its principal uses have been the degradation of alkaloids 158-162 and the cleavage of an alkyl group from N,N-dialkylanilines. 163,164 The von Braun cleavage is discussed in detail in Chapter 4.

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142 Collie and Sehryver, J. Chem. Soc., 57, 767 (1890).
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150 Wedekind, Ber., 35, 766 (1902).

153 Meyer and Lecco, Ann., 180, 173 (1876).

154 Wedekind and Pasehke, Ber., 43, 1303 (1910).

155 von Braun, Ber., 33, 1438 (1900); Scholl and Norr, ibid., 33, 1550 (1900).

155 Elderfield and Hageman, J. Org. Chem., 14, 605 (1949).

ur von Braun and co-workers, Ber., 33, 2728, 2734 (1900); 35, 1279 (1902); 40, 3933 (1907); 41, 2100, 2113 (1908); 42, 2035, 2219 (1909); 43, 1353, 3209 (1910); 44, 1252, (1911); 47, 3023 (1914); 51, 96, 255 (1918); 55, 3803 (1922); 56, 1840, 2165 (1923); 63, 2407 (1930); 70, 1241 (1937); Ann., 445, 201 (1925); 449, 249 (1926); 490, 189 (1931); 507, 1 (1933).

us Mossler, Monatsh., 31, 1 (1910).

us von Braun, Ber., 47, 2312 (1914); 49, 2624 (1916).

162 Speyer and Sarre, Ber., 57, 1427 (1924). 151 Speyer and Rosenfeld, Ber., 58, 1125 (1925).

12 Leuchs and Overberg, Ber., 65, 961 (1932); 66, 79 (1933).

103 von Braun, Ber., 37, 2670 (1904); 40, 3914 (1907); 41, 2165 (1908).

14 Sachs and Weigert, Ber., 40, 4356 (1907).

¹⁵¹ Miehler and Gradmann, Ber., 10, 2078 (1877). 152 Marquardt, Ber., 19, 1027 (1886).

Replacement of Amine by Hydrogen (Emde Reduction)

$$[RN(CH_3)_3]^+X^- + 2(H) \rightarrow RH + N(CH_3)_3 \cdot HX$$

 $RN(CH_3)_2 + H_2 \rightarrow RH + NH(CH_3)_2$

Quaternary salts may be reduced either by means of sodium amalgam (Emde reduction) 165-168 or lithium aluminum hydride, 168a or catalytically; 169,170 tertiary amines are subject to catalytic reduction only. 154, 169-175 Many of these reductions are discussed in Chapter 5. Phenolic Mannich bases can also be reduced by means of sodium methoxide.175a

RELATED SYNTHETIC PROCESSES

Carbon-carbon alkylation by amine replacement is of particular value when a labile amino compound is more readily accessible as a starting material than is the corresponding halide or conjugated unsaturated compound. The following section is intended to place the reactions that have been discussed in perspective relative to other methods that result in the formation of similar products or are formally related to the amine replacement reactions. For obvious reasons, no attempt has been made to cover these aspects of synthetic organic chemistry in a detailed or exhaustive manner.

Carbon-Carbon Alkylations by Halogen Replacement

Some of the most familiar and important methods for the formation of carbon-carbon bonds involve replacement of the halogen atom of an

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165 Emde, Ber., 42, 2590 (1909).
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¹⁶⁵ Emde and Kull, Arch. Pharm., 272, 469 (1934).

¹⁶⁷ Groenewoud and Robinson, J. Chem. Soc., 1934, 1692. 168 von Braun and co-workers, Ber., 49, 501, 1283, 2613 (1916); 50, 50 (1917); 55, 3803

^{(1922); 56, 1570 (1923).} 1634 Kenner and Murray, J. Chem. Soc., 1950, 406.

¹⁶⁹ Emde, Helv. Chim. Acta, 15, 1330 (1932).

¹⁷⁰ Emde and Kull, Arch. Pharm., 274, 173 (1936).

DITKOIET, Ber., 75, 429 (1942).

173 Baltzly and Buck, J. Am. Chem. Soc., 65, 1984 (1943); Baltzly and Russel, J. Am.

¹⁷³ Caldwell and Thompson, J. Am. Chem. Soc., 61, 765 (1939). Chem. Soc., 72, 3410 (1950).

¹⁷⁴ Bachman and Levine, J. Am. Chem. Soc., 69, 2341 (1947). Bachman and Levine, J. Am. Chem. Soc., 70, 686 (1948); Carlin and Landerl, J. Am. 13 May and Mosettig, J. Am. Chem. Soc., 70, 686 (1948); Carlin and Landerl, J. Am.

May and Mosettig, J. Am. Chem. Soc., 10, 3252 (1950); Karrman and Bladh. Chem. Soc., 72, 2762 (1950); Reeve and Sadle, ibid., 72, 3252 (1950); Karrman and Bladh. Acta Chem. Scand., 4, 1541 (1950) [C. A., 45, 7092 (1951)].

cta Chem. Scand., 4, 1541 (1950) [C. A., 40, 1052 (1957), 1942, 682; Rapoport, King, and 1752 Cornforth, Cornforth, and Robinson, J. Chem. Soc., 1942, 682; Rapoport, King, and 1752 (1957) Lavigne, J. Am. Chem. Soc., 73, 2718 (1951).

alkyl halide. 176 Some of the more important of these methods are the following:

Friedel-Crafts Reaction. 177

$$RX + ArH \xrightarrow{AlCl_3, etc.} RAr + HX$$

Reaction with Organometallic Compounds. 178

$$RX + MR' \rightarrow RR' + MX$$

Alkylation of Active Methyl and Methylene Compounds. 179

$$\begin{array}{c} RX + Na - \begin{matrix} Z \\ C - R' \\ Y \end{matrix} \rightarrow \begin{array}{c} Z \\ R - \begin{matrix} C - R' + NaX \\ Y \end{matrix}$$

A particularly interesting example of this type of reaction represents a new route to cyclohexenones such as may be prepared by use of ketonic Mannich bases.²⁹

Mannich bases. H₅C₂O₂C
$$H_5$$
CCl=CHCH₂Cl+ H_5 C₂O₂C H_5 CCl=CHCH₂Cl+ H_5 C₂O₂C H_5 CH₃CCl=CHCH₂Cl+ H_5 CO₂R H_2 SO₄ H_2 SO₄ H_2 SO₄ H_3 CC H_4 CH H_5 CH

Replacement by Cyanide. 180

$$RX + MCN \rightarrow RCN + MX$$

¹⁷⁶ Weygand, Organic Preparations, pp. 353-403, Interscience Publishers, New York, 1945.

Price in Adams, *Organic Reactions*, Vol. III, p. 1, John Wiley & Sons, 1946.
 See Ref. 176, pp. 355-358.

¹⁷⁹ See Ref. 176, pp. 355–358.

¹⁸⁰ See Ref. 176, p. 367.

Carbon-Carbon Alkylations by Oxygen Replacement 181

Friedel-Crafts Reaction.

$$ROR' + ArH \rightarrow R-Ar + HOR'$$

This reaction is not of general applicability.

Alkylations with Ethylene Oxide.

O
$$CO_2R'$$
 CO_2R' CO_2R'
 CH_2 — CH_2 + Na — C — R'' $NaOCH_2CH_2C$ — R''
 Z

and/or

 CH_2 — O
 CH_2 — O
 CH_2 — O
 CH_2 — C

Alkylations with o-Hydroxybenzyl Alcohols.

$$n \longrightarrow^{\text{CH}_2\text{OH}} \longrightarrow \left[\longrightarrow^{\text{OH}}_{\text{CH}_2} \longrightarrow^{\text{CH}_2} \right] n$$
(as well as para and crosslinked polymers)

Alkylation with Diethyl Methoxymethylmalonate. 182, 183

Alkylation with Diethyl Mothers
$$+ CH_3OCH_2CH(CO_2C_2H_5)_2 \rightarrow \\ CO_2C_2H_5 + CH_3OCH_2CH(CO_2C_2H_5)_2 + CH_3OH \\ CO_2C_2H_5 + CH_$$

Carbon-Carbon Alkylations with Diazoacetic Ester. 184

Carbon-Carbon Alkylations with 2
$$\frac{CH_2CO_2C_2H_5}{N} + N_2CHCO_2C_2H_5 \rightarrow \frac{N}{CH_3}$$

¹⁸¹ See Ref. 176, pp. 404-414.

¹⁵² Fischer and Nenitzescu, Ann., 443, 113 (1925).

¹³³ Maurer and Moser, Z. physiol. Chem., 161, 131 (1926).

¹⁵⁴ Piccini, Gazz. chim. ital., 29, 363 (1899).

Carbon-Carbon Alkylation by Sulfur Replacement 185

Coupling of Active Hydrogen Compounds by Condensation with Carbonyl Compounds

One-Step Condensations. A. Formation of Symmetrical Products.

$$\begin{array}{c} R & R \\ | & | \\ C = 0 + HA \rightarrow A - C - A + H_2 0 \\ | & | \\ R & R \end{array}$$

This process is useful in the formation of symmetrical compounds, except that it cannot be used when the hydrogen in H—A and the α -hydrogens in RCOR are of comparable activity. Phenols, malonic esters, β -keto esters, α -cyano esters and secondary amines are among the types of compounds that will undergo symmetrical coupling of the type shown above. A familiar example is the synthesis of DDT.

B. Formation of Unsymmetrical Products.

$$AH + C = O + HB \rightarrow A - C - B + H_2O$$

When the active hydrogen compounds to be coupled are markedly different in structure and activity, good yields of unsymmetrical products

¹⁸⁵ Cardwell, J. Chem. Soc., 1949, 715.

may be obtained. The halo-alkylation 186 and amino-alkylation (Mannich) reactions 1 (p. 103) of active hydrogen compounds are well-known examples of unsymmetrical coupling reactions. The reaction of Nmethylolamides with aromatic compounds 6,187 is a less familiar example.

$$C_5H_5CONH_2 + CH_2O \rightarrow C_6H_5CONHCH_2OH$$

$$C_6H_6CONHCH_2OH + OCH_3 \rightarrow OCH_3$$

The cyanomethylation of indole 188,189 is one of the few examples in which two different carbanion-forming substances can be coupled by means of formaldehyde to yield unsymmetrical products.

The first step in Two-Step Condensation-Addition Reactions. reactions of this type is the familiar Perkin-Claisen-Knoevenagel reaction; 190,191 the second step consists in addition of an active hydrogen compound to a conjugated unsaturated system (Michael reaction).192 An example is the synthesis of phenylsuccinic acid. 193

$$\begin{array}{c} C_{6}H_{5}CHO + CH_{2}(CN)CO_{2}H \rightarrow \\ C_{5}H_{5}CH = C(CN)CO_{2}H \xrightarrow{C_{2}H_{5}OH} C_{6}H_{5}CH = C(CN)CO_{2}C_{2}H_{5} \\ C_{6}H_{5}CH = C(CN)CO_{2}C_{2}H_{5} + HCN \rightarrow \\ C_{6}H_{5}CHCHCO_{2}C_{2}H_{5} \xrightarrow{and \\ decarboxylation} C_{6}H_{5}CHCH_{2}CO_{2}H \end{array}$$

This process can be employed to advantage when the conjugated unsaturated compound is easily prepared and stable enough to be isolated and purified, and it is of principal value when the carbonyl compound which serves as a coupling agent is some material other than formaldehyde. In many syntheses, an active hydrogen compound can be added to a conjugated unsaturated compound, such as acrolein or acrylonitrile, 194 which is more easily prepared in some other way. In

¹⁸⁸ Fuson and McKeever in Adams, Organic Reactions, Vol. I, p. 63, John Wiley & Sons,

¹⁸⁷ Einhorn, Ann., 343, 207 (1905); 361, 113 (1908); Downes and Lions, J. Am. Chem.

¹⁸³ Bauer and Andersag, U. S. pat. 2,222,344 [C. A., 35, 1807 (1941)]. Soc., 72, 3053 (1950).

¹⁵⁷ Sankyo, Jap. pat. 161,544 [C. A., 43, 2236 (1949)].

Johnson in Adams, Organic Reactions, Vol. I, p. 210, John Wiley & Sons, 1942.

Jonnson in Adams, Organic Iteactoria, An Advanced Treatise, Vol. I, pp. 122 Allen and Blatt in Gilman, Organic Chemistry. An Advanced Treatise, Vol. I, pp. 672-688, John Wiley & Sons, New York, 1944.

¹³³ Lapworth and Baker, Org. Syntheses Coll. Vol. 1, 181, 451 (1941). 134 Bruson in Adams, Organic Reactions, Vol. V, p. 79, John Wiley & Sons, 1949.

such syntheses the relationship of synthetic methods is only formal, though the products obtained are structurally similar to those formed by the two-step condensation-addition process outlined above.

CHOICE OF EXPERIMENTAL CONDITIONS

Choice of Reactants

Carbon-carbon alkylations of hydrogen cyanide and active methyl or methylene compounds by Mannieh bases are part of a two-step process for coupling active hydrogen compounds by means of formaldehyde with the loss of water. It is often theoretically possible to form ZCH₂Z'

$$ZH + CH2O + HN(R)2 \rightarrow ZCH2N(R)2 + H2O$$
$$ZCH2N(R)2 + HZ' \rightarrow ZCH2Z' + HN(R)2$$

by alkylation of ZH with the Mannich base of HZ'. It is apparent, then, that it may at times be necessary to decide which active hydrogen compound should be converted to its Mannich base and which should be reserved as the compound to be alkylated if best yields are to be obtained.

If the formation of γ -ketonitriles and aryl- or indole-acetonitriles or acetic acids is desired, there seems to be no alternative to the use of ketonic, phenolic, or indole Mannieh bases. At any rate, the use of α -aminoacetonitriles as alkylating agents is seldom feasible. However, when the desired process is the coupling of two active hydrogen compounds, neither of which is hydrogen eyanide, there is often a choice of which one to employ as a Mannich base and which as the reagent to be alkylated.

The following points should be considered.

1. The Mannich reaction takes place readily with compounds containing even only moderately active methyl or methylene groups.

- 2. Only compounds containing highly active methylene groups are easily alkylated by means of Mannich bases. Compounds that contain only moderately active methylene groups, such as simple ketones, usually require the presence of strong bases such as sodium amide capable of converting them to enolates if they are to be alkylated by Mannich bases.
- 3. Only those tertiary amines that can form conjugated unsaturated systems by amine elimination are suitable for use as alkylating agents (p. 126).
- 4. Only those quaternary ammonium salts that can suffer amine elimination or that possess allylic systems are suitable for use as alkylating agents (p. 104).

In the cases under consideration, then, it is desirable to convert to its Mannich base the active methyl or methylene compound possessing the least acidic hydrogen atoms, provided, of course, that this Mannich base can undergo amine elimination or possesses an allylic system; and to use the appropriate active methylene compound possessing the most acidic hydrogen atom (only one such active hydrogen atom is necessary) as the reagent to be alkylated.

Mannich bases that can suffer amine elimination possess active hydrogen atoms and could, conceivably, be subject to alkylation. Intermolecular self-alkylation of the Mannich base (p. 103) should be most prominent in alkylations of compounds containing hydrogen atoms whose acidity is similar to or less than that of the active hydrogen atoms of the Mannich base. It is probable that this accounts for the facts that phenolic Mannich bases give only diarylmethanes in attempted base-catalyzed alkylations of active methylene compounds, and large amounts of diarylmethanes in their reactions with hydrogen cyanide, and that low yields of the desired alkylation product are usually obtained when ketones are alkylated by quaternary salts of ketonic Mannich bases. In the latter case, formylation of the ketone prior to alkylation may result in an improved yield (p. 114).

advantages, since their quaternary salts are often more soluble in inert solvents than the corresponding halides.

Ease of Purification of the Mannich Bases or Their Salts. Many of the simpler Mannich bases of ketones may be purified by distillation; high temperatures are to be avoided, because amine elimination may occur. It is advantageous to use the relatively low-boiling dimethylamino Mannich bases.

Ketonic Mannich bases are best stored as their hydrochlorides, in which form they are usually isolated. Dimethylamino, piperidino, and morpholino Mannich base hydrochlorides are particularly easily crystallized. The dimethylamino and morpholino Mannich base hydrochlorides are often appreciably more hygroscopic than the piperidine derivatives.

The piperidino and morpholino Mannieh bases of phenols are generally crystalline and stable, whereas a number of the dimethylamino, diethylamino and, especially, dibutylamino Mannieh bases of phenols are thick liquids which are not always distillable. Most of the Mannieh bases of indole are crystalline and stable.

Quaternary salts of Mannich bases are often too unstable to permit long storage. Indeed, quaternary salts of some phenolic Mannich bases decompose at room temperature or lower at rates that preclude their isolation. Wilds and Shunk ²⁶ have shown the necessity of using pure quaternary salts of ketonic Mannich bases in alkylations of active methylene compounds if good yields of pure products are to be obtained. Piperidino, dimethylamino, and, especially, morpholino Mannieh bases form easily crystallizable quaternary salts.

Inertness of the Amine Undergoing Replacement. Derivatives of aniline would generally be expected to be unsuitable for use in carbon-carbon alkylations by amine replacement because of the ease with which nuclear substitution in the aromatic amine could occur.

Volatility of the Amine Undergoing Replacement. The elimination of amines from Mannich bases is reversible. ^{67,109} If the secondary amine formed during the reaction is not removed, it could compete for the conjugated unsaturated compound with the substance to be alkylated. As quaternary salts of Mannich bases can undergo facile amine exchange reactions with tertiary amines, ^{65,c,84} amine elimination from such salts is probably reversible too, and removal of the tertiary amine by volatilization would seem to be desirable also. Trimethylamine (b.p. 3.5°), dimethylamine (b.p. 7.4°) and diethylamine (b.p. 55°) are readily distilled from the reaction mixture during the reaction when the solventials, for example, ethanol. Piperidine (b.p. 106°) and morpholine (b.p. 126–130°) could be removed in this manner only when higher boiling

solvents, such as hexanol, toluene, xylene, dibutyl ether, or Diethyl Carbitol, are used. One of the most convenient methods for following the course of an amine replacement reaction is observation of the evolution of a volatile amine.

Choice of Solvents, Operating Temperatures, etc.

The choice of solvents, reaction temperatures, reaction times, and apparatus to be used in amine replacement reactions varies according to the nature of the reaction and will be considered in more detail in the following sections. A few general remarks can be made at this point, however.

Mannich bases and their salts seem to be sensitive to air oxidation in alkaline reaction media and at temperatures required for some of the reactions. Although it is not invariably necessary to employ an inert atmosphere, such as nitrogen, in these reactions, it would seem to be generally desirable. A slow nitrogen stream also serves to sweep volatile amines out of the reaction mixture, thus making it somewhat easier to follow the reaction, which may be assumed to be completed when amine evolution (detected by odor or by moist red litmus paper) ceases.

Experimental Conditions for Particular Types of Carbon-Carbon Alkylations

Replacement of Amino Groups by Cyanide. Use of Ketonic Mannich Bases. The method of Knott ¹³ seems to be generally applicable for the formation of γ -ketonitriles from the hydrochlorides of dimethylamino Mannich bases of aryl methyl ketones (see preparation of β -benzoyl-propionitrile, LXIV, p. 155) and requires no comment. It is possible propionitrile, this procedure may be required if other types of Mannich bases are employed.

$C_6H_5COCH_2CH_2CN$ LXIV

Use of Mannich Bases of Indoles and Phenols. A solution of the Mannich base and an excess (100-500%) of sodium cyanide in aqueous ethanol is heated under nitrogen with reflux until the evolution of secondary amine and ammonia is complete or greatly reduced (36-80 secondary amine and ammonia is complete or aryl-acetic acid formed may be conhours) (Hood). The indole- or aryl-acetic acid formed may be conhours with the corresponding acetamide, and, when phenolic taminated with the corresponding acetamide, and, when phenolic Mannich bases are used, with diarylmethanes or phenol-formaldehyde resins are genresins. The diarylmethanes and phenol-formaldehyde resins are genreally insoluble in sodium carbonate; their removal is illustrated in the

preparation of 3-indoleacetic acid, p. 155, and 2-hydroxy-1-naphthaleneacetic acid, p. 156.

The conversion of Mannich bases of phenols and indoles to nitriles by reaction with hydrogen eyanide in benzene at 150° in an autoclave has been described only in the patent literature.¹²

Use of Quaternary Salts of Benzylamines. The use of these salts is of little synthetic importance in the benzene series since the corresponding benzyl chlorides are often easily prepared by chloromethylation. The quaternary salts of furfurylamines (XII, R = II) and especially 5-methylfurfurylamines (XII, R = CII₃) may prove useful, for the amines are more easily prepared and handled than the corresponding halides and may give rise to different products (p. 107).

The method consists in either distilling an aqueous solution of the quaternary salt and alkali cyanide at atmospheric pressure to remove all the water or simply mixing the dry quaternary salt with dry sodium cyanide and then carefully heating the residue or mixture in vacuum to a temperature of 150-200° so that the nitrile distils as it is formed. Overheating should be avoided, since the reaction may become quite violent. The nitrile is usually contaminated with the tertiary amine corresponding to the quaternary salt. In another modification of the technique, an aqueous paste of the quaternary salt and the cyanide is heated to about 200°, and the nitrile formed is swept out with superheated steam at the same temperature.¹⁰

Alkylation of Active Methyl and Methylene Compounds

Although conditions for the alkylation of active methyl and methylene compounds by means of Mannich bases are in general similar, rather wide variations in procedure have been employed. The following generalizations can be made, however.

An inert atmosphere is generally employed; apparently Mannich bases or intermediates in the alkylation reactions (such as vinyl ketones) are sensitive to oxygen, yielding tars or colored products as a result of oxidation or free-radical catalyzed polymerization.

As previously pointed out, ionization of the compound to be alkylated is necessary if the reaction is to occur. This ionization may be caused by the basic character of the Mannich base itself (as in alkylations of ethyl nitroacetate ¹⁹ or tricarbethoxymethane ¹⁷) or by added sodium hydroxide, sodium ethoxide (as in alkylations of derivatives of malonic ester or of β -keto esters), or sodium amide (as in alkylations of ketones). It seems best to use a base that is no stronger than necessary, if multiple alkylations and other condensation reactions are to be avoided.

Only catalytic amounts of added base are required when Mannich bases are employed as alkylating agents; the base may be added to a mixture of the Mannich base and the substance to be alkylated. Alkylations under these conditions are often very slow. Some alkylations proceed just as well or even better in the absence of base.²¹⁶

When quaternary ammonium salts are the alkylating agents, an equivalent amount of base is necessary since it is consumed during the In practice, the sodium enolate of the active methylene compound is first formed, and the quaternary salt is then added to the reaction mixture. Alternatively, a quaternizing agent such as methyl iodide can be added to a mixture of the Mannich base and the sodium enolate of the active methylene compound to be alkylated. Alkylations of this type have only rarely been carried out without solvent; occasionally an excess of the active methylene compound to be alkylated is the solvent. It is obvious that solvents possessing hydrogen atoms more acidic than those of the substance to be alkylated are unsuitable for these reactions, since they would destroy the enolate. Thus, except in alkylations of such strongly acidic substances as diethyl cyanomalonate, water seems to have a deleterious effect (see, however, ref. 213). Absolute ethanol has been widely used as a solvent in alkylations of malonic esters and β -keto esters. The sodium enolates of ketones are generally formed by reaction with sodium amide in ether, pyridine, or benzene; and the quaternary salt alkylating agent, suspended in the same solvents or dissolved in an alcohol, is then added. In the alkylation of 1-methyl-5-methoxy-2-tetralone (LXV) with diethylaminobutanone 35 (p. 160), potassium ethoxide was satisfactory as a condensing agent; in this

instance the methiodide of the Mannich base (formed on the walls of the reaction vessel) and a benzene solution of the ketone were brought together first and the base was then added in ethanolic solution. This technique gave an unusually high yield (71%) of the alkylation product. In alkylations of malonic ester derivatives by gramine (VIIa) (in the presence of powdered sodium hydroxide), toluene or xylene (which presence of powdered sodium hydroxide). Polyfunctional dissolve both reactants) has been used successfully. Polyfunctional high-boiling ethers, such as Diethyl Carbitol, are good solvents for the sodio derivatives of active methylene compounds and seem to dissolve

appreciable amounts of some quaternary ammonium salts; such ethers may prove to be useful solvents in alkylations of sodium enolates of active methylene compounds by means of quaternary salts.

EXPERIMENTAL PROCEDURES

The formulas of certain of the substances described in the following preparations are given herewith for purposes of reference.

$$\begin{array}{c} \text{CH}_2\text{CH}(\text{CO}_2\text{H})_2 \\ \text{CH}_3 \\ \text{LXXVIII} \end{array}$$

 β -Benzoylpropionitrile (LXIV). To a mixture of 213.5 g. (1.0 mole) of β -dimethylaminopropiophenone hydrochloride 195 (XXV, $\hat{R} = CH_3$) and 130 g. (2.0 moles) of potassium cyanide in a 5-l. flask is added 2.6 l. of boiling water. The mixture, consisting of an aqueous and an oily layer, is heated under reflux for thirty minutes. Part of the dimethylamine which is evolved distils and may be collected in dilute hydrochloric acid. When the mixture is chilled, the oil solidifies and crystals separate from the water layer. β -Benzoylpropionitrile (105 g., 67%) is separated by filtration and is crystallized from a benzene-light petroleum mixture, forming almost colorless blades, m.p. 76°.

3-Indoleacetic Acid (XX) and 3-Indoleacetamide. 16* 25.0 g. (0.144 mole) of gramine (3-dimethylaminomethylindole, VIIa),196 35.2 g. (0.717 mole) of sodium cyanide, 280 ml. of 95% ethanol, and 70 ml. of water is boiled under reflux for eighty hours. To the cooled reaction mixture is added 350 ml. of water. The solution is treated with Norit, filtered, concentrated under reduced pressure until all the ethanol has been removed, cooled to 5°, and filtered. The solid on the funnel is recrystallized from ethanol and ether to give 5.0 g. (20%) of 3-indoleacetamide, m.p. 149-151°.

The reaction mixture, after removal of the amide by filtration, is concentrated under reduced pressure to a volume of about 300 ml. and cooled to 10°. Dropwise addition of cold, concentrated hydrochloric acid (Hood!) to the vigorously stirred solution causes precipitation of crude, slightly pink 3-indoleacetic acid. The crude material is filtered and dried at 70°; yield, 20.0 g. (79%) of material melting at 158-161°.

^{*} Because an alkali cyanide is used, this preparation should be run in a well-ventilated hood. The waste liquors must be handled and disposed of with care, since they contain considerable amounts of cyanide.

¹⁹⁵ Maxwell, Org. Syntheses, 23, 30 (1943).

¹⁹⁸ Kühn and Stein, Ber., 70, 567 (1937).

The crude product can be recrystallized from ethylene dichle ide containing a small amount of ethanol to give 17.4 g. (69%) of pure 3-indole-acetic acid, melting at 167–168°.

A solution of 1.0 g. of 3-indoleacetamide, 1.2 g. of sodium hydroxide, and 8 ml. of water is boiled for four hours. The cooled solution (5°) is treated with Norit, filtered, and made strongly acid (about pH 1.5) with concentrated hydrochloric acid. The acid which precipitates is collected by filtration and dried at 70°. The yield of 3-indoleacetic acid melting at 167–168° is 0.95 g. (95%). The over-all yield of 3-indoleacetic acid from gramine is 88%.

2-Hydroxy-1-naphthaleneacetic Acid (XVIII). A solution of 4.2 g. (0.02 mole) of 1-dimethylaminomethyl-2-hydroxynaphthalene 197,193 (IV, R = H) and 2.09 g. (0.04 mole) of sodium cyanide in 60 ml. of 50% ethanol is heated under reflux in a nitrogen atmosphere for thirty-six hours, at the end of which time the evolution of dimethylamine and ammonia is complete. The flask is cooled to room temperature under a slow stream of nitrogen, and the yellow solution is poured without delay into 100 ml. of water. Dry Ice (100 g.) is added to the solution in small portions (hydrogen cyanide is evolved in this process). The white precipitate which forms when the solution is saturated with carbon dioxide is removed by filtration and washed with water. This material is crude di-(2-hydroxy-1-naphthyl)methane (XIX).

To the filtrate is added 50 g. of ice, and then slowly and with stirring, under a good hood, 50 ml. of concentrated hydrochloric acid, whereupon glistening platelets of 2-hydroxy-1-naphthaleneacetic acid separate. This material is collected by filtration, washed with 10% hydrochloric acid and then with water. There is obtained 1.90 g. (47%) of air-dried product melting at 146.5°. Often the material has a steel-blue color; the color can be removed by dissolving the acid in aqueous sodium carbonate, filtering, and reprecipitating with acid.

1-β-Indolyl-2-nitrobutane (LXVI). A mixture of 10 g. (0.058 mole) of gramine ¹⁹⁶ (VIIa), 50 ml. of redistilled 1-nitropropane, and 2.6 g. of solid sodium hydroxide is heated under reflux for six to eight hours or until amine evolution ceases. The solution is cooled and acidified with 50 ml. of 10% aqueous acetic acid and is then diluted with 200 ml. of ether. It is then washed with four 75-ml. portions of water, shaken with Norit, and filtered. The solvents are distilled at room temperature under

^{*} Because an alkali cyanide is used, this preparation should be run in a well-ventilated hood. The waste liquors must be handled and disposed of with care, since they contain the formula of cyanide.

Décombe, Compt. rend., 197, 258 (1933).
 Ger. pat. 89,979 [Frdl., 4, 98 (1899)].

vacuum, leaving a viscous oil. Distillation of the oil at 157°/0.2 mm. furnishes 10.6-11.9 g. (82-95%) of $1-\beta$ -indolyl-2-nitrobutane, m.p. 90-91°.

Ethyl Skatylnitroacetate (XXIII).19 In a 250-ml. flask, fitted with a stirrer, a thermometer, a nitrogen inlet, and a condenser, is placed 8.66 g. (0.05 mole) of gramine 196 (VIIa), 13.3 g. (0.10 mole) of ethyl nitroacetate, 199,200 and 50 ml. of dry xylene. A slow stream of nitrogen is passed through the vigorously stirred mixture, and the temperature is raised to 90-100° and held there for five hours. (About one-half the calculated amount of dimethylamine may be collected in a trap through which the exit gases pass.) The hot solution is filtered, and the xylene is distilled under reduced pressure. The residual gum is taken up in chloroform, and the solution is extracted with two 50-ml. portions of 10%hydrochloric acid solution and then washed with water until neutral. The chloroform solution is dried over magnesium sulfate, the chloroform is removed by distillation at 20-30 mm. and the excess ethyl nitroacetate is distilled at 1 mm. The oil that remains is dissolved in chloroform, and the solution is extracted with successive portions of 5% sodium hydroxide solution until the oil that separates on acidification of a test portion is negligible in amount. The combined basic extracts are acidified with 10% hydrochloric acid, the temperature of the mixture being kept below 20°, and then extracted with chloroform. The chloroform solution is dried and concentrated; the residual oil crystallizes readily. The yield of ethyl skatylnitroacetate is 11.8 g. (90%). The melting point of a sample recrystallized from benzene-petroleum ether is 62.0-62.8°.

2-(2'-Nitroethyl)cyclohexanone (LXVII). A mixture of 15 g. (0.097 mole) of 2-dimethylaminomethylcyclohexanone 201 (XVI) and 9.2 g. (0.151 mole) of nitromethane is heated on a steam bath; 27 ml. of a 10% solution of sodium methoxide in methanol is added in one portion with vigorous stirring. As soon as the reaction product becomes solid. the solution is diluted with 20 ml. of methanol and stirred without further heating until the evolution of dimethylamine is complete. The sodium salt of the product is dissolved in water; the solution is cooled in an ice-salt bath and acidified with acetic acid. The red-brown oil that separates is taken up in several portions of ether, and the combined ethereal solutions are washed with water and dried over sodium sulfate. The ether is distilled, and the residual oil (12 g., 72%) is distilled at 160°/14 mm. for purification.

¹⁹⁰ Steinkopf, Ber., 42, 3925 (1909); Ann., 434, 21 (1923).

²⁰⁰ Feuer, Hass, and Warren, J. Am. Chem. Soc., 71, 3078 (1949).

²⁰¹ Mannich and Braun, Ber., 53, 1874 (1920).

2-Keto-3-carbethoxy-9-hydroxydecalin (LXVIII).²⁴ A mixture of 16 g. of 2-dimethylaminomethylcyclohexanone ²⁰¹ (XVI) and 15 g. of ethyl acetoacetate is treated on the first, third, fifth, and seventh days of standing with a solution of 0.1 g. of sodium in 3 ml. of absolute ethanol. The solution turns yellow, then red. After fourteen days the reaction is complete. (Further addition of sodium ethoxide only lowers the yield; it is not advantageous to add the alkoxide in one portion.) After the first four or five days of standing, crystals form in the solution and rapidly increase in bulk. The crystal mass is filtered from the green fluorescent liquid and washed with dilute hydrochloric acid and a little water. After recrystallization from ethanol and drying, the fine white needles weigh 18 g. (73%) and melt at 146°.

2-Keto-3-carboxy- $\Delta^{1,9}$ -octalin (LXIX).²⁴ Six grams of the ethyl ester LXVIII and 1.7 g. of potassium hydroxide are dissolved in 40 ml. of cold water. The solution is allowed to stand four days and is then treated with sulfuric acid until acid to Congo Red; a powdery mass precipitates. The aromatic odor of this material is probably due to the presence of 2-keto- $\Delta^{1,9}$ -octalin (LXX). The acid is best purified by dissolving it in cold sodium carbonate solution and reprecipitating with hydrochloric acid. The material is obtained in good yield and melts at 95° with liquefaction and loss of carbon dioxide.

2-Keto- $\Delta^{1,9}$ -octalin (LXX).²⁴ When the keto acid LXIX is melted, 2-keto- $\Delta^{1,9}$ -octalin boiling at 140-141°/14 mm. is formed in yields of about 75%.

2-y-Ketobutyl-2-carbomethoxy-1-keto-1,2,3,4-tetrahydrophenanthrene (LXXI).26 The sodium enolate of 2-carbomethoxy-1-ketotetrahydrophenanthrene 202 (LXXII) is prepared by heating 2.07 g. of the keto ester with a solution of 0.19 g. of sodium in 10 ml. of dry thiophenefree benzene. The mixture, containing the insoluble sodium salt, is then cooled in an ice bath, and the methiodide from 2.5 g. of redistilled 1-diethylamino-3-butanone 26 (XXIV, $R=C_2H_5$) is added in 10 ml. of methanol. The sodium salt slowly dissolves, and, after four hours at room temperature, another crystalline precipitate separates. mixture is refluxed for one hour. The clear solution is then diluted with water and extracted twice with benzene. After the solution has been washed with water, dilute acid, and water, the benzene is evaporated and the residue is crystallized from ethyl acetate to give 2.31 g. of cubic prisms, m.p. 141-143°. A second crop (0.18 g., m.p. 130-140°) is a mixture of prisms and needles which can be separated by adding petroleum ether, suspending the lighter needles by swirling, and decanting

²⁰² Bachmann and Wilds, J. Am. Chem. Soc., 62, 2084 (1940).

them with the liquid. Recrystallization of the residue affords 0.12 g. more of the prisms, m.p. 139-142°, making the total yield 92%. When further purified by distillation at 0.5 mm. and recrystallization from ethyl acetate, the material melts at 145-145.3°.

Cyclization of the Tricyclic Keto Ester LXXI to the Tetracyclic Keto Ester LXXIII. One gram of the keto ester (product melting above 139° is suitable) and a solution of 1 g. of sodium in 100 ml. of anhydrous methanol are heated at reflux for two hours under nitrogen. After the solution has been cooled, water is added and the mixture is extracted with two portions of benzene. The benzene solution is washed with water, evaporated, and the residue crystallized from ethyl acetate to yield 0.52 g. of yellow needles, m.p. 174–176°. A second crop (0.23 g., m.p. 160–175°) brings the total to 79%. After the second crystallization from ethyl acetate (Norit), the product LXXIII melts at 178.5–179.5°.

The keto ester can be hydrolyzed and decarboxylated to the tetracyclic ketone LXXIV in 52% yield with aqueous methanolic potassium hydroxide.

Cyclization of the Tricyclic Keto Ester LXXI to Form the Tetracyclic Ketone LXXIV. (a) A mixture of 500 ml. of methanol, 5 ml. of 45% potassium hydroxide, and 0.8 g. of the keto ester LXXI is heated under reflux under a nitrogen atmosphere for twenty hours. The solution is diluted, and the product is extracted with three portions of warm benzene. The extract is washed with water and dilute hydrochloric acid and then concentrated. The first crop of 0.40 g. of yellow plates melts at 182–185°, and the second crop at 176–183°. The total yield is 90%. A sample purified for analysis by evaporative distillation at 0.5 mm. and recrystallized from benzene forms colorless plates, m.p. 188–188.5°.

(b) When 0.5 g. of the diketo ester LXXI is refluxed with 25 ml. of acetic acid and 5 ml. of hydrochloric acid under nitrogen, the product LXXIV obtained by dilution and extraction with benzene weighs 0.32 g. (84%) and melts at 185-187°.

2-Keto-10-methyl- $\Delta^{1,9}$ -octalin (LXXV).²⁵ A mixture of 33 g. of 2-methylcyclohexanone, 6.1 g. of powdered sodium amide, and 50 ml. of dry ether is stirred for four hours in a stream of dry nitrogen at room temperature. A solution of 43 g. of 1-diethylamino-3-butanone methiodide ²⁶ in 20 ml. of absolute ethanol is then slowly added, and after four hours the solution is heated under reflux for two hours. Dilute hydrochloric acid and ether are added, and the ethereal solution is separated, dried, and distilled, giving 9.3 g. (38%) of 2-keto-10-methyl- $\Delta^{1,9}$ -octalin, b.p. 143–145°/16 mm.

7-Keto-1-methoxy-13-methyl-5,6,7,9,10,13-hexalydrophenanthrene (LXXVI).35 Fifteen grams of methyl iodide is added in portions during one-half hour to 15.05 g. of 1-diethylamino-3-butanone 26 (XXIV, R = C₂H₅), which is swirled gently in a 1-1, flask cooled in ice. The swirling is regulated to obtain the crystalline methiodide as an even coating on the walls of the flask. When no more liquid remains, the flask is kept in ice for one-half hour and then under the tap for 45 minutes. A solution of 20.0 g. of 1-methyl-5-methoxy-2-tetralone 35 (LXV) in 100 ml. of dry, thiophene-free benzene is added, the air is expelled from the flask by dry nitrogen, and a solution of potassium ethoxide prepared from 6.5 g. of potassium and 100 ml. of dry ethanol is added with ice cooling during five minutes. Swirling is continued until all the methiodide has dissolved (about 30 minutes) and has been replaced by a precipitate of potassium iodide. After the mixture has been kept in ice for another hour, it is boiled gently for twenty-five minutes. An excess of 2 N sulfurie acid is then added, and the nitrogen stream is stopped. After addition of enough water to dissolve the potassium sulfate, the benzene layer is separated and the aqueous layer extracted twice with ether. The combined organic extracts are washed with water, dried with a little magnesium sulfate, and evaporated. The residue is distilled to yield 23.2 g. of material boiling up to 180°/1 mm. The distillate is warmed until fluid, and ether is added gradually until the total weight is 40 g. Crystallization begins at once and is allowed to proceed at 0° overnight. The product is collected and washed with chilled ether; it weighs 17.0 g. and melts at 115-117°. Fractional distillation of the mother liquors affords an additional gram of material, making the total yield 71%. The process has been carried out successfully on four times the above scale.

Diethyl Skatylacetamidomalonate (XLIV). To a boiling mixture of 1.2 l. of toluene (or xylene) and 17 g. of powdered sodium hydroxide contained in a 5-l. three-necked flask fitted with a mechanical stirrer, a condenser, and a nitrogen inlet tube are added 250 g. (1.43 moles) of gramine ¹⁹⁶ (VIIa) and 311 g. (1.43 moles) of diethyl acetamidomalonate (XLI). Refluxing and rapid stirring are continued for five hours while a rapid stream of nitrogen is passed through the reaction mixture. The evolution of dimethylamine, which is very rapid at the beginning, almost ceases at the end of the heating period.

The reaction mixture is filtered through a preheated funnel, and the filtrate is held at 5° for several hours to aid crystallization. The product is collected by filtration and washed with cold toluene followed by petroleum ether. The dried product (446 g., 90%) melts at 158-159°

and can be converted without further purification to (+,-)-tryptophan by the method of Snyder and Smith.⁴⁵

Ethyl β-(2-Pyrrolyl)-α-cyano-α-acetamidopropionate (LXXVII).⁴³ In a flask fitted with a stirrer, a condenser, and a dropping funnel are placed 100 ml. of absolute ethanol and 1.72 g. of clean sodium. After all the sodium has dissolved, 12.7 g. of ethyl acetamidocyanoacetate ²⁰³ (XLV) and then 9.3 g. of 2-dimethylaminomethylpyrrole ²⁰⁴ are added. While the flask is cooled in an ice bath and the mixture is stirred, 15.8 g. of dimethyl sulfate is added dropwise at such a rate that the temperature does not exceed 35°. After the addition is complete, the mixture is stirred for one hour and allowed to stand at room temperature for about eight hours. The ethanol is evaporated under reduced pressure, and the residue is diluted with 200 ml. of water and chilled. Nearly white crystals (17 g., 90%) separate. They are purified by dissolving in acetone, treating with charcoal, filtering, diluting with water, and chilling for several hours. The white plates which form melt at 122°.

This material can be hydrolyzed and decarboxylated in one step by treatment with hot sodium hydroxide.

1-Methylskatylmalonic Acid (LXXVIII).¹⁷ To a solution of 0.23 g. (0.01 gram atom) of sodium in 30 ml. of absolute ethanol are added 4.65 g. (0.02 mole) of tricarbethoxymethane 205 and 3.3 g. (0.01 mole) of 1-methylgramine methiodide (IX).9 The mixture is refluxed for one and one-half hours under a current of nitrogen; there is a vigorous evolution of trimethylamine. While refluxing is continued, 10 ml. of 40% aqueous sodium hydroxide is added, followed, after ten minutes, by 10 ml. of water. Trimethylamine evolution resumes for some time. After about two hours, heating is discontinued and the solution is concentrated under reduced pressure, extracted twice with ether, acidified with concentrated hydrochloric acid, and chilled. The brown crystals which separate are collected and dissolved in 15 ml. of a saturated solution of sodium carbonate and 25 ml. of water. The solution is boiled with Norit, filtered with suction, acidified with concentrated hydrochloric acid and chilled. The light pink crystals after thorough washing with ice water and drying weigh 1.55 g. (62%) and melt at 171-172° (dec.).

A 34.5% yield can be obtained when the alkylation is carried out in aqueous medium.

²⁰³ Tullar, U. S. pat. 2,393,723 [C. A., 40, 2465 (1946)].

²⁰⁴ Herz, Dittmer, and Cristol, J. Am. Chem. Soc., 69, 1698 (1947).

²⁰⁵ Lund and Voigt, Org. Syntheses Coll. Vol. 2, 594 (1943).

TABULAR SURVEY OF ALKYLATION PRODUCTS

In Tables I-X are summarized carbon-carbon alkylations with amines and ammonium salts reported prior to January 1, 1951. Some more recent references have been included in the text but not in the tables. Certain references may have been overlooked, since there is no sure way of locating the alkylation reactions in the literature.

Yields are given as stated in the original literature. A dash indicates

that no yield is reported.

Table I is concerned with carbon-carbon alkylations with aliphatic and aromatic tetraalkylammonium salts other than phenolic Mannich bases which are listed in Table II. Table III contains alkylations with heterocyclic Mannich bases by the Mannich method (p. 113), Table IV similar alkylations by the Robinson method (p. 113), and Table V analogous reactions by the Albertson method (p. 118). Table VI is concerned with alkylations with ketonic Mannich bases; in Table VII are listed alkylations of alkali metal cyanides with the hydrochlorides, and in Table VIII various alkylations with the quaternary salts of such bases. Table IX contains a survey of an alkylation of indole with diethyl piperidinomethylformamidomalonate under a variety of conditions, and in Table X are listed some recently reported alkylations with Mannich bases of nitro compounds.

Within each table, the reactions are arranged in order of complexity of the alkylating group, and, for the same alkylating group, in order of the compounds alkylated, cyanides being listed first, nitro compounds next, then esters and ketones, and last organometallic compounds.

TABLE I CARBON ALKYLATIONS WITH ALIPHATIC AND AROMATIC TRIALKYLAMINES AND TETRAALKYLAMMONIUM SALTS

				Viola	Refer-
Quaternary Salt	Compound Alkylated	Solvent	Product	%	ence
Tetramethylammonium cyanide		None	Acetonitrile and methylcarbyl- amine	_	3
Dimethylethylanilinium iodide	Potaesium cyanide	None	Acetonitrile		4
Tetramethylammonium ethexide	Diethyl malonate	Ethanol	Diethyl met hylmalonate	58	38
Tetramethylammonium ohloride	9-Fluoryllithium	None	9-Methylfluorene	-	38
Benzyldimethylanilinium chloride	Potassium cyanide	None	Benzyl cyanide		4
Benzyldimethylanilinium chloride	Sodium cyanide	Water	Alkylation failed		7
Benzyltrimethylammo- nium iodide	2-Nitropropane sodium salt	Ethylene glycol	Benzaldehyde	Low	206
Benzyltrimethylammo- nium bromide	Diethyl eodiomalonate	Di-n-buty ether	l Diethyl benzylmalonate	77	7
Benzyltriethylammonium ìodide	Diethyl eodiomalonate	Di-n-buty ether	l Diethyl henzylmalonate	63	7
Benzyldimethylanilinium chloride	Diethyl sodiomalonate	Ethanol	Diethyl benzylmalonate	38	7
Benzyldimethylanilinium chloride	Diethyl sodiomalonate	None	Diethyl henzylmalonate	73-79	7
Benzyldimethylanilinium chloride	Diethyl sodiomalonate	Ethanol (auto- clave)	Diethyl henzylmalonate	32–36	7
Benzyldimethylanilinium ethoxide	Diethyl sodiomalonate	Ethanol	Diethyl benzylmalonate	51	7
Benzylmethylpiper- idinium chloride	Diethyl sodiomalonate	Ethanol	Diethyl benzylmalonate	5	7
Banzylmethylpiper- idinium chloride	Diethyl sodiomalonate	Ethanol (auto- clave)	Diethyl benzylmalonate	20-26	7
Benzylmethylpiper- idinium iodide	Diethyl eodiomalonate	Ethanol (auto- clave)	Diethyl henzylmalonate	22-36	7
Benzylmethylpiper- idinium jodide	Diethyl sodiomalonate	Di-n-hutyl ether	Diethyl henzylmalonate	43	7
Benzyltrimethylammo-		None	Hydrocinnamonitrile	-	
nium cyanoacetate			Dibenzylacetonitrile	_	39
Benzyldimethylamine	Methyl cyanoscetate	None	Dibenzylmethylamine Hydrocinnamonitrile	15	28
Бенгунишеску вание	менци сувновсеные	24046	Dibenzylacetonitrile	19	
			Dibenzylmethylamine	23	39
Benzyldimethylanilinium chloride	Ethyl sodioacetoacetate	Ethanol	Ethyl benzylacetoacetate	60	7
Benzyldimethylamine	Tricarbethoxymethane	None	Hydrocinnamio acid Dibenzylacetic acid	39 19	39
Benzyltrimethylammo- nium hromide	Phenyllithium	Ether	1,1,2-Triphenylethane		61
Benzyltrimethylammo- nium iodide	Phenylmagnesium hromide	Di-n-butyl ether	Biphenyl	25	62
Note: References 206-22	9 are listed on p. 197.				

TABLE I-Continued

CARBON ALKYLATIONS WITH ALIPHATIC AND AROMATIC TRIALKYLAMINES AND TETRAALKYLAMMONIUM SALTS

				Yield	Refer-
Quaternary Salt	Compound Alkylated	Solvent	Product	%	ence
Benzylpyridinium chlo- ride	Phenylmagnesium bromide	Di-n-butyl ether	Biphenyl	4	62
p-Nitrobenzyltrimethyl- ammonium iodide	2-Nitropropane sodium salt	Ethanol	p-O ₂ NC ₆ H ₄ CH ₂ C(CH ₁) ₂ NO ₂	63	206
(+)-α-Phenylethyltrimeth- ylammonium iodide	Sodium cyanide	None	Styrene		2
(+)-α-Phenylethyltrimeth- ylammonium iadide	Diethyl sodiomalonate	Diethyl Carbitol	(+, -)-a-Phenylethylmalonic ester	18	2
 Dimethylaminomethyl- 2-methoxynaphthalene methiodide 	Sodium cyanide	None	2-Methoxy-1-naphthylaceto- nitrile	44	6
1-Dimethylaminomethyl- 2-methoxynaphthalene methiodide	Diethyl sodiomalonate	Diethyl Carbitol	CH ₂ CH(CO ₂ H) ₂ OCH ₃	61	6
C&H&CH(N)	Benzylmagnesium chloride	Ether	C6H5CHCC6H5	19	63
CeH2CH(N)CH3	$egin{align*} \mathbf{Benzylmagnesium} & \mathbf{chloride} \ 2 \end{aligned}$	Benzene	C6H6CH N CH3	15	63
Note: References 206-22	9 are listed on p. 197.		CH ₂ C ₆ H ₅		

TABLE II

CARBON-CARBON ALKYLATIONS WITH 0-HYDROXYBENZYLAMINES

Substituted o-Hydroxybenzylamine	Compound Alkylated	Solvent; Temperature	Product	Yield %	Refer ence
	-	-	Poduci		
2-Dimethylaminomethyl- phenol	Phenylmagnesium bromide	Di-n-butyl ether; reflux	_	0	62
2-Dimethylaminomethyl- 6-methoxyphenol	Sodium cyanide	90% ethylene glycol, 10% water; reflux	2-Hydroxy-3-methoxy phenylacetic acid	4	150
2-Dimethylaminomethyl- 6-methoxyphenol	Ethyl cyanoaco- tate	Excess ethyl cyanoace- tate; 190°	Resins and N,N-dimethyl- cyanoacetamide		150
2-Dimethylaminomethyl- 4-methylphenol	Sodium cyanide	50% ethanol; 150°	2-Hydroxy-5-methylphenyl- acetic acid	_	12
2-Dimethylaminomethyl- 4-methyl-6-hromophenol	Sodium cyanide	50% ethanol; 150°	2-Hydroxy-3-hromo-5-methyl- phenylacetic acid		12
2-Dimethylaminomethyl- 4-allyl-6-methoxyphenol	Sodium cyanide	50% ethanol; 150°	2-Hydroxy-3-methoxy-5-allyl- phenylacetic acid	-	12
2-Diethylaminomethyl-4-allyl- 6-methoxyphenol	Sodium cyanide	50% ethanol; 150°	2-Hydroxy-3-methoxy-5-allyl- phenylacetic acid	-	12
1-Dimethylaminomethyl- 2-hydroxynaphthalene	Sodium cyanide	50% ethanol; reflux	2-Hydroxy-1-naphthalene- acetic acid	47	12, 15
			Di-2-hydroxy-1-naphthyl- methane	20	15
I-Dimethylaminomethyl- 2-hydroxynaphthalene	Hydrogen cyanide	Benzene; 150°	2-Hydroxy-1-naphthalene- acetonitrile		12
1-Morpholinomethyl-2-hy- droxynaphtbalene	Dihenzoyl- methane	Ethanolic HCl; reflux	CH ₂ CH(COC ₆ H ₅) ₂ OH	53	23
1,5-Bis(dimethylaminomethyl)- 2,6-dihydroxynaphthalene	Sodium cyanide	50% ethanol;	2,6-Dihydroxynaphthalene- 1.5-diacetic acid		12
5-Dimethylaminomathyl- 6-hydroxyquinoline	Sodium cyanide	50% ethanol; 150°	6-Hydroxyquinoline-5-acetic acid	_	12

Note: References 206-229 are listed on p. 197.

TABLE III

Carion Alkylations with Heferocyclic Compounds Containing a Dialkylaminomethyl Group

Amine 2-Dimethylamino- methylpyrrole	Compound Alkylated Diethyl malonate	Solvent; Catalyst Toluene; NaOH	Product CH_2 CH_2 $C = C - CH_2$	Yield %	Reference ence 43
2-Dimethylamino- methylpyrrole	Diethyl acetamido- malona <i>to</i>	Toluene or xylene; NaOH	$O = C - N$ C_{12} $C_{2}C_{2}H_{6}$	70-80	43
2-Dimethylamíno- methylpyrrole	Diethyl benzamido- malonate	Xylene; NaOH	O=C—C NHCOCH ₃ NHCOCH ₃ NHCOCH ₃ O=C—C	38	43a
Gramine (3-di- methylamino-	Sodium cyanide	Ethanol-water; none	$NHCOC_6H_6$ Indole-3-acetic acid Indole-3-acetic acid	20	16
methylindole) Gramine	Sodium cyanide	Ethanol-water;	Indole-3-acetic acid	Quant.	12
Gramine	Hydrogen eyanide	Benzene; none	Indole-3-acetonitrile	1	12

		C	ARBO	N AI	KYI	ATI	ONS WI	rh A	MINE	3		167	
12	12	18	18	18	18	18	18 19 20	7	17	52a	41	41	
1	١	20	20	82-95	82		80 90 97	92	29	86	06	48	
Indole-3-acetic acid	Indole-3-acetonitrile	Diskatylnitromethane *	1-Nitro-1-skatylethane *	- 1-Nitro-1-skatylpropane *	- 2-Nitro-2-skatylpropane *	Alkylation failed	Ethyl diskatylnitroacetate * Ethyl skatylnitroacetate * Diethyl skatylnitromalonate *	Diethyl skatylmalonate *	Skatylmalonic acid *	Diethyl skatylformamidomalonate *	Diethyl skatylacetamidomalonate *	Diethyl skatylacetamidomalonate *	
Ethanol-water;	none Renzene: none	Excess nitro- methane:	NaOH Excess nitroeth-	ane; NaOH Excess 1-nitropro-	Excess 2-nitropro-	pane; NaOH Excess 2-nitro-	1-butanol Ethanol; NaOH Xylene; none	Excess diethyl	malonate; Na Excess tricar- bethoxymeth-	ane; none Toluene; NaOH	Xylene or tolu-	ene; NaOH Pyridine; none	
; ;	Sodium cyanide	Hydrogen cyamae Nitromethane	Nitroethane	1-Nitropropane	9. Nitropropage	2-Nitro-1-butanol	Ethyl nitroacetate Ethyl nitroacetate	Diethyl nitromalonate Diethyl malonate	Tricarbethoxymethane	Diethyl formamido-	malonate Diethyl acetamide	malonate Diethyl acetamido- malonate	
	3-Diethylamino- methylindole	3-Piperidino- methylindole Gramine		Gramine Gramine		Gramine	Gramine Gramine Gramine	Gramine	Gramine	3-Diethylamino-	methylindole Gramine	Gramine	

TABLE III—Continued

Heterocyclic Compounds Containing a Dialkylaminomethyl Group

Dofor	relei-	ence	41	41	41	41	41	62	102a			17	17, 39
1.1	Y 1eld	%	55	54	98	64	10	လ	46				10
CONTRACTOR		Product	Diethyl skatylacetamidomalonate *	Diethyl skatylacetamidomalonate *	Diethyl skatylacetamidomalonate *	Diethyl skatylacetamidomalonate *	Diethyl skatylphthalimidomalonate st	3-Benzylindole	(CII ₂ NC	C _c H _s	Alkylation failed	Di-(1-methyl-3-indolyl)methane 1-Methylskatylmalonic acid *
ROCYCLIC COMPOUNDS		Solvent;	Catalyst Pyridine; NaOH	No solvent; none	Toluene; NaOH	Toluene; NaOH	Toluene; NaOH	Di-n-butyl ether;	none	Xylene; Na		Ethanol-water;	none Exeess methyl cyanoaeetate;
ATTENDED OF THE HETEROCYCLIC COMPOUNDS CONTINUED	WINDLAM INC.	Compound	Alkylated	Dietayi acetamide malonate Diethyl gestamide-	Dietiryi accoming malonate Diethyl gestamido-	Dietiyi acceanido malonate Diethyl costamido-	Dietiyi aceraniya malonate Diethyl phthalimido-	malonate Phenylmagnesium	bromide	NCOC6Hs CN		Sodium cyanide	Methyl cyanoacetate
,000	CARBO		Amine	Gramine	Gramine	3-Diethylamino- methylindole	3-Piperidino- methylindole Gramine		Gramme	Gramine		1-Methylgramine	1-Methylgramine

17	17	17	200	707	53	27	55	102a		
15	15 9	4 8 15	- 1	18	1	92	61	69		
Di-(1-methyl-3-indolyl)methane 1-Methylskatylmalonic acid *	1-Methylskatylmalonic acid * 3'-(1-Methyl-3-indolyl)propionic acid	1-Methylskatylacetamidomalonic acid * 1-Methyl-N-acetyltryptophan Di-(1-methyl-3-indolyl)methane	Alkylation falled	4-Chloroindole-3-acetamide	Diethyl 5-bromoskatylacetamido-	malonace Ethyl 6-methylskatylacetamidocyano-	acetate " Diethyl 2-carbethoxyskatylacetamido- malonate *	NON NO	CO ₂ C ₂ H ₅ N	ц П
Excess ethyl cyanoacetate;	Na Excess tricarbeth- oxymethane;	none Excess diethyl acetamidomal- onate; Na	Di- n -butyl ether;	Ethanol-water;	Xylene; NaOH	Xylene; NaOH	Xylene; NaOH	Xylene; Na		
Ethyl cyanoacetate	Tricarbethoxymethane	Diethyl acetamido- malonate	Methylmagnesium iodide	Potassium cyanide	Diethyl acetamido-	malonate Ethyl acetamidocyano-	acetate Diethyl acetamidomalo- nate	The Cooper	CN	
1-Methylgramine	1-Methylgramine	1-Methylgramine	1-Methylgramine	4-Chlorogramine	5-Bromogramine	6-Methylgramine	3-Diethylamino- methyl-2-car-	bethoxylindole 3 Dimethylamino- methyl-2-car-	bethoxyindole	

Note: References 206-229 are listed on p. 197.

TABLE III-Continued

CARBON ALKYLATIONS WITH HETEROCYCLIC COMPOUNDS CONTAINING A DIALKYLAMINOMETHYL GROUP

Refer-	ence	208	ដ	500	200	
Yield	૾ૢૺ	‡ 0g	ı			
	Product	CH2C(CO3C2H2)2	2-Phenylindole-3-acetic acid	Alkylation failed	Alkylation failed	
Solvent;	Catalyst	Xylene; NaOH	Ethanol-water; none	Xylene; NaOH	Toluene; none	
Compound	Alkylated	Diethyl acetamidomalonate	Sodium cyanide	3-Piperidinomethyl- Diethyl nitromalonate indazole	3-Piperidinomethyl- Diethyl acetamidomalo- indazole nate	
	Amine	3-Dimethylamino- methylindole- 2-carboxy- piperidide	2-Phenyl-3-di- methylamino- methylindole	3-Piperidinomethyl- indazole	3-Piperidinomethyl- indazole	, 4 , 1

Note: References 206-229 are listed on p. 197. * The skatyl group is

† The yield was based on the acid obtained by hydrolysis of the amide. ‡ Over-all yield from indole-2-carboxypiperidide.

TABLE IV

Carbon Alkylations with Salis of Heterocyclic Compounds Containing a Trialkylaminomethyl Group

Refer- ence	10	10	43		62	2	r- £	- C - T	50 45, 46	
Yield $\%$	22-32 4-5	37	53			46	85	8	50 63-70 45, 46	
Product	2-Furylacetonitrile 5-Methvl-2-furonitrile	5-Methyl-2-furylacetonitrile	, CH	O=C—C NHCOCH3	Di-n-butyl ether Alkylation failed	Indoleacetonitrile	Skatylmalonie aeid * Diethyl skatylmolonate *	Diethyl skatyleyanoacetate * Ethyl skatylecetoacetate *	Diethyl skatylacetamido-	malonate *
Solvent	Water or none	Water	Dioxane		Di-n-butyl ether	Water	Di-n-butyl ether Not reported	Di-n-butyl ether Di-n-butyl ether Ethanol	Dioxane	
Compound Alkylated	Sodium cyanide	Sodium cyanide	Diethyl sodioacetamido- malonate		Methylmagnesium iodide	Potassium silver cyanide	Diethyl sodiomalonate Diethyl sodiomalonate	Ethyl sodioeyanoacetate Ethyl sodioacetoacetate Ethyl sodioacetoacetate	Diethyl sodioacetamido-	בוניים סווים ביי
Quaternary Salt	Furfuryltrimethylammonium iodide	2-Dimethylaminomethyl- 5-methylfuran methiodide	2-Dimethylaminomethyl- pyrrole methiodide		2-Dimethylaminomethyl- pyrrole methiodide	Gramine (3-dimethylamino- methylindole) methiodide	Gramine methiodide Gramine ethiodide	Gramine methiodide Gramine methiodide Gramine ethiodide	Gramine methiodide	

Note: References 206-229 are listed on p. 197.

TABLE IV-Continued

•	Refer-	0000
THYL GROUP	Yield	Š
MONIM PART TARE	gnds Containing a litabathering	
TABLATION AT MARKET	Yield Refer-	ARBON MURITARIONS HELD
	(٧

Cinnon ALKYLATIONS WILL CALL	WITH CALCALL			1	
TOWN TOWN		,	Product	%	cnec
Quaternary Salt	Compound Alkylated	Solvent Ethanol	Diethyl skatylacetamido-	١	46
Gramine methiodide	Diethyl sodioacetamido- malonate		malonato * Dicthyl skatylacetamido-	١	47
Gramine ethiodide	Diethyl sodioacetamido- malonate	, 1	malonate * Diethyl skatylbenzamido-	١	47
Gramino ethiodide	Dicthyl sodiobenzamido- malonate	Total other	malonate * Diethyl skatylphthalimido-	١	46
Gramine methiodide	Diethyl sodiophthalimido- malonate	Di-Ar-Dutyi cuitor	malonate *	8	ę
Gramine methiodide	Methylmagnesium iodide	Di-a-butyl culer	sym-Di-3-indolylethane	16	29
Gramine methiodide	Phenylmagnesium bromide	Di-n-butyl ether	3-Benzylindole sym-Di-3-indolylethane	24	62
Gramine methiodide	Benzylmagnesium chloride	Di-n-butyl ether	3-(Phenethyl)indole 1-Methylindole-3-acetonitrile	60-64	, •
1-Mothylgramine methiodide	Sodium eyanide	17 24 001	1,3-Dimethyl-2-cyanoindole	22	17
1-Methylgramine methiodide Diethyl sodiomalonate	Diethyl sodiomalonate	Ethanol or ex- eess diethyl	1-Methylskatylmalome acid		
1-Methylgramine methiodide Ethyl sodiocyanoacetate	Ethyl sodiocyanoacetate	malonate Execss ethyl evanoacetate	1-Methylskatylmalonic acid *·†	17	17
4. H.	Tricorpothoxymethane	Ethanol	1-Methylskatylmalonic acid *,†	63	17
1-Methylgramine methiodide		Water	1-Methylskatylmalonic aeid *.†	35	17
1-Methylgramine methiodide	ricarbellosymetras sodium enolate				

1-Methylgramine methiodide	Ä	Water or ethanol	Water or ethanol 1-Methylskatylmalonic aeid *'†	51	17
1-Methylgramine methiodide	苺	Ethanol	Ethyl 1-methylskatylaeet- amidoevanoaeetate *	69	48
1-Methylgramine methiodide 1-Methylindole	cyamorceave 1-Methylindole	Aqueous ethanol	Di-(1-methyl-3-indolyl)- methane	49	17
1-Methylgramine methiodide 1-Methylgramine methiodide	Methylmagnesium iodide Phenylmagnesium bromide	Di-n-butyl ether Di-n-butyl ether	1-Methyl-3-ethyl indole 1-Methyl-3-benzyl indole	44	62 62
1-Methyl-5-methoxygramine methiodide	Diethyl sodioacetamido- malonate	Ethanol	Diethyl 1-methyl-5-methoxy-skatylacetamidomalonate *	98	210
3-Piperidinomethylindazole methiodide	Diethyl sodioacetamido- malonate	Ethanol	CH2C(CO2C2H5)2 N NHCOCH3	35 †	209
3-Dimethylaminomethyl- indazole methiodide	Ethyl sodioacetamidocy- anoacetate	Ethanol	CN CH2CC2C2H6 CH2CCC2H6	i	500
Note: References 206-229 are listed on p. 197.	e listed on p. 197.		H		111 /

Note: References 200-229 are listed on p. 150.

† The acid was obtained by hydrolysis of the primary alkylation product.

FPOUNDS CONTAINING A DIALEXLAMINOMETHYL GROUP USING DIMETHYL Š

Refer-	43, 43a 43a 43a	43	43a	43a	43a	43a	43	43a	43		
Yield	33, 44 Low 38	30	8	55	63	27	94	84	90		
TEROCYCLIC CONPOUNDS CONTAINING AGENT SULFATE OR ETHYL IODIDE AS A QUATERNIZING AGENT	Product 2-C4H4NCH2CH(CO2C2H6)2* (2-C4H4NCH2)2C(CO2C2H6)2*	Ethanol; dimethyl (2-C,H4,NCH2)2C(COZOZOZOZ) sulfate ' '', C,H,NCH2)2C(CN)CO2C2H5*	Ethanol; dimethy! (2-C,H4)\(CH2CH(CO2C,H6)\)2.	(2-C,H,NCH ₂ C(CO ₂ C,H ₆)* 2-C,H,NCH ₂ C(CO ₂ C,H ₆)* CH ₃	Takend: dimethyl 2-C,H4NCH2C(CO2C2H6)?*	sulfate C ₆ H ₅ Sulfate 2-C ₄ H ₄ NCH ₂ C(CO ₂ C ₂ H ₅)2*	sulfate OCOCH3	Ethanoi; dimethyl 2.C.H4NCH2C(CO2C2H5);*	NHCOC ₆ H ₆ CN	2-C4H4NCH2CC02C2H8*	NHCOCH3
ounds Contamily yl Iodide as a Qu	Solvent; Quater- nizing Agent Ethanol; dimethyl sulfate	Ethanol; dimethyl sulfate	Ethanol; dimetayi sulfate	Ethanol; dimethyl sulfate	Tetranol dimethyl	sulfate	Sulfate			Ethanol; dimetary sulfate	
CARBON ALKYLATIONS WITH HETEROCYCIJO COMPOUNDS CONTALIZACION CARBON ALKYLATIONS WITH HETEROCYCIJO COMPOUNDS CONTALIZACION ALKYLATION ALKYLATIONS WITH HETEROCYCIJO COMPOUNDS CONTALIZACION ALKYLATION ALKYLAT	Compound Alkylated Diethyl sodiomalonate	Diethyl sodiomalonate	Ethyl sodiocyanoacetate	Tricarbethoxymethane sodium enolate Diethyl sodiomethyl-	malonace	Dicthyl sodiophenyl- malonate	Diethyl sodioncetoxy- malonate	Diethyl sodioacetamido- malonate	Diethyl sodiobenzamido- malonate	Ethyl sodioacetamido- cyanoacetate	
CARBON ALKYLATIONS V	Amine 9. Dimethylaminomethyl-	pyrrole pyrrole pyrrole	Pyrrolo (2 moles) 2-Dimethylaminomethyl-	2. Dimethylaminomethylprinor pyrrolo	pyrrolo	2.Dimethylaminomethylpyrrole	2-Dimethylaminomethylpyrrole	2-Dimethylaminomethyl- pyrrole	2-Dimethylaminomethyl- pyrrole	2-Dimethylaminomethyl-	P31161

	CA	RBON ALI	CYLATIONS W	ITH AMINES	175
43	44	44	#	44	8 40, 50 40, 8
Low	100	1	48.5	20	50 † 65, 82, 86, 72, 82
Ethanol; dimethyl 2-C4H4NCH2C(CO2C2H6)2* sulfate NCO	(H,C2O2C)2CCH2 CH2C(CO2C2H5)2 CH3CONH H NHCOCH3	CH CCH ₂ C(CO ₂ C ₂ H ₈) ₂ N S NHCOCH ₃	H3CC——CCH2CH(CO2C2Hs)2	H ₃ CC——CCH ₂ C(CO ₂ C ₂ H ₅) ₂ N S NHCOCH ₃	HCOCH3 Indole-3-acetonitrile Diethyl skatylacetamidomalonate † Diethyl skatylacetamidomalonate †
Ethanol; dimethyl 2 sulfate	Ethanol; dimethyl sulfate	Ethanol; diethyl sulfate and so- dium ethoxide	Ethanol; dimethyl sulfate	Ethanol; dimethyl sulfate	Aqueous ethanol; dimethyl sulfate Ethanol; ethyl iodide Ethanol; dimethyl sulfate (1 mole)
Diethyl sodiophthali- midomalonate	Diethyl sodioacetamido- malonate	Diethyl sodioacetamido- malonate	Dicthyl sodiomalonate	Diethyl sodioacetamido- malonate	Potassium cyanide Diethyl sodioacetamido- malonate Diethyl sodioacetamido- malonate
2-Dimethylaminomethyl- pyrrole	2,5-Bis(piperidinomethyl)- pyrrole	2-Acetamido-5-dimethyl- aminomethylthiazole bydrochloride	2-Acetamido-4-methyl- 5-dimethylaminomethyl- thiazole hydrochloride	2-Acetamido-4-methyl- 5-dimethylaminomethyl- thiazole	Gramine (3-dimethylam- inomethylindole) Gramine Gramine

51

23

Diethyl 6-methylskatylacetamido-

5

93

Ethanol; dimethyl Diethyl 7-methylskatylacetamido-

malonate ‡ malonate ‡

Ethanol; dimethyl

sulfate sulfate

Ethanol; dimethyl

Diethyl sodioacetamido-Diethyl sodioacetamido-

malonate

malonate malonate malonate ‡

sulfate

Diethyl sodioaeetamido-

malonate

7-Methylgramine

6-Methylgramine

5-Methylgramine

TABLE V-Continued

5 27 Refer-210 $\overline{2}$ 20 ence 22 යි 8 40 49 CARRON ALKYLATIONS WITH HETEROCYCLIC COMPOUNDS CONTAINING A DIALKYLAMINOMETHYL GROUP USING DIMETHYL 33 83 8 91 ١ 83 20 86 Ethyl skatylsuccinimidocyanoacetate ‡ Ethyl skatylphthalimidoacetoacetate ‡ Diethyl skatylphthalimidomalonate ‡ Ethanol; dimethyl Ethyl skatylacetamidoeyanoacetate ‡ Ethyl skatylacetamidoacetoacetate ‡ Diethyl 5-methoxyskatylacetamido-malonate ‡ Diethyl skatylbenzamidomalonate ‡ Diethyl 5-methylskatylaeetamido-Diethyl skatylbenzamidomalonate ‡ Ethanol; dimethyl Diethyl skatylacetamidomalonate † sulfate (1.65 Diethyl skatylacetamidomalonate ‡ Diethyl 4-methylskatylacetamido-Diethyl 2-methylskatylacetamido-Product malonate ‡ malonate ‡ Ethanol; dimethyl Ethanol; dimethyl sulfate Ethanol; dimethyl Ethanol; dimethyl Ethanol; dimethyl Ethanol; dimethyl Solvent; Quater-Ethanol; ethyl Ethanol; ethyl Ethanol; ethyl Ethanol; ethyl sulfate sulfate iodide iodide sulfate sulfate mole) Diethyl sodiobenzamido-Diethyl sodioacetamido-Diethyl sodioacetamido-Diethyl sodioacetamido-Diethyl sodiobenzamido-Diethyl sodioacetamido-Ethyl sodiophthalimido-Ethyl sodiosuccinimido-Diethyl sodioacetamido-Diethyl sodiophthalim-Compound Alkylated Ethyl sodioacetamido-Ethyl sodioacetamidocyanoacetate acetoacetate cyanoacetate acetoacetate idomalonate malonate malonate malonate malonate malonate 3-Piperidinomethylindole 3-Diethylaminomethyl-5-Methoxygramine 2-Methylgramine 4-Methylgramine Amine Gramine Gramine Gramine Gramine Gramine Gramine Gramine

	CARBON	ALKYLATI	ONS WITH AMIN	(ES 17	'
42 52 $39a$	39a	39a	39a		
65 87 43	44	33	21 28		
Ethanol; dimethyl Diethyl 4-cyanoskatylmalonate ‡ sulfate Ethanol; dimethyl Ethyl 5-methylskatylacetamidocyano- sulfate acetate ‡ acetate ‡ CH2CHCO2C2H5 sulfate	$R = CO_2C_2H_5$ $[ethy] \qquad \qquad CH_2CH_2CHCO_2C_2H_5$	R = COC	methyl CH ₂ CH ₂ CH ₂ CH ₃ N(CH ₃) ₂ (7) $CH_2 CH_2 CH(CO_2 C_2 H_6)_2(7)$	nitrile,	
Ethanol; dimethyl sulfate Ethanol; dimethyl sulfate Ethanol; dimethyl sulfate	Ethanol; dimethy ^l sulfate	Ethanol; dimethyl sulfate	Ethanol; dimethyl sulfate	Irolysis of the	
Diethyl sodiomalonate Ethyl sodioacetamido- cyanoacetate Diethyl sodiomalonate	Ethyl sodioacetoacetate	Ethyl sodiobenzoyl- acetate	Diethyl sodiomalonate	Note: References 206-229 are listed on p. 197. * 2-C ₄ H ₄ N represents MH	NH.
4-Cyanogramine 3-Dicthylaminomethyl- 5-methylindole 2-β-Dimethylaminocthyl- quinoline	2-6-Dimethylaminoethyl- quinolinc	2-6-Dimethylaminoethyl- quinoline	CHICHE-NICHS)2l2	Note: References 206–2: * 2-C,H,N represents † The yield was based o † The skatyl group is	

TABLE VI

WITH B-AMINOKETONES

Refer-	cnce 21	56	1	56 56	211		30	211	212		212
	k	1	18	1.1	1		42	40	1		40
	Cyclized Product	1,3-Cyclohexanedione	3-Methyl-4-carbethoxy-	2-cyclone-2-cyclobezen-1-one 3-Methyl-2-cyclobezen-1-one 4-Carbethoxy-4-isopropyl- 3-methyl-2-cyclobezen-	$\frac{1-\text{one}}{\text{CO}_2}$ C ₂ H ₅		2-Keto-10-phenyl-∆¹, 9-octalin	$\begin{array}{c} co_2 c_2 H_6 \\ \\ \\ \\ \end{array}$	OCH3	١	C ₆ H ₆ CH ₂ CH ₂ CO ₂ H
S WITH P-rames	Yield Cyclizing Agent	NaOCAH.		ı	53 Mg + I2		1	1		18	1
ALKYLATIONS OF ACTIVE METHYLENE COMPOUNDS WITH PARTICULAR	Yi Simple Alkylation Product		CH ₃ COCH ₂ CH ₂ CH(CO ₂ C ₂ H ₆) ₂	l	1	CO ₂ C ₂ H ₅ CH ₂ CH ₂ COCH ₃			I	HO2C(CH2)2CO(CH2)2CH(CO2C2H5)2	1
ALKYLATIONS	Active Methylene Compound (Con-	densieg Agent) Nitromethane	(NnOCH3) Diethyl malonate	(NaOC2115) Ethyl acetoacetate (NaOC2H5)	Ethyl isopropyl- acetoacetate (NaOC ₂ H ₅)	2-Carbethoryey- cleheranone (NaOC2Hs)		2-Phenylcyclo- hexanone (NaNH2)	2-Carhethoxycy- clehexanone (NaOC2Hs)	Dicthyl malonate	(NaOC2Hs) Phenylacetone (KOC4H9-t)
		β-Anineketone	none 1-Dimethylamino-3-buta-	none 1.Dimethylamino-3.buta- oene	1-Dimethylamino-3-buta- nene	1-Diethylnmiao-3-buta- nene		1-Dinethylamino-3-buta- nooc	1-Diethylnmino-3-penta- nooc	4-Keto-6-dimethylamino-	eaproio acid hydrochlo- ride 4-Keto-G-dimethylamino- caproio acid hydrochlo- rido

21	21	21	213	214	21		21	25	21	22	215	77
53	į	1	91	1	1 1	1	1	18	I	1	1	73
2-Phenylpyrrolidine	1	!	O=C $O=C$	3-Phenyl-6-carbethoxy-2-cy- clohexen-1-one	3-khenyi-z-cyclobexen-i-one	í	1	CO ₂ C ₂ H ₅	CH ₃		0	2-Keto-3-carbethoxy-9-hy- droxydecalin
$Z_{n}(Hg)+HCl$	I	1	HCl + CH₃CO₂H	I	í	í	1	1	i	$KOH + C_2H_5OH;$ then $(CH_3CO)_2O$	I	I
23	1	1	52	∞	1	l	1	ĺ	72	81	19	1
γ -Nitrobutyrophenone	(C,H,COCH,CH,),CHNO,	(C6H5COCH2CH2)3CNO2	CH ₂ CH ₂ COC ₆ H ₅	Ethyl acetoacetate C ₆ H ₅ COCH ₂ CH ₂ CH(CO ₂ C ₂ H ₅)COCH ₃ (NaOC ₂ H ₅)	γ -Nitro-4-methory $butyrophenone$	7-Nitro-3,4-dimethoxybutyrophenone	CH ₃ O COCH ₂ CH ₂ CHNO ₂		2-p-Nitroethylcyclohexan-1-one	CH2CH(CO2C2H5)2 =0	CH ₂ CH(CO ₂ C ₂ H ₆) ₂	1
Nitromethane (KOH + CH,OH)	Nitromethane	(KOH + CH ₃ OH)	Cyclohexanone (NaOH + H2O + C ₂ H ₅ OH)	Ethyl acetoacetate (NaOC ₂ H ₅)	Nitromethane	Nitromethane	(NaOCH ₃)	Ethyl acetoacetate (NaOC2Hs)	Nitromethane (NaOCH ₃)	Diethyl malonate (NaOC ₂ H ₅)	Diethyl malonate (None)	Ethyl acetoacetate (NaOC ₂ H ₅) 9 are listed on p. 197
β-Dimethylaminopropio-	g-Dimethylaminopropio-	phenone	β-Dimethylaminopropio- phenono hydrochloride	β-Dimethylaminopropio- plienone	β-Dimethylamino-4-meth- oxvpronionhenone	\$-Dimethylamino-	5,4-dimethoxypropio- phenone	$CH_2N(C_2H_6)_2$ CH_3	2-Dimethylaminomethyl- cyclohexanone	2-Dimethylaminomethyl- cyclobexanone	2-Dimethylaminomethyl- cyclohexanone	2-Dimethylaminomethyl- Ethyl acetoacetate cyclohexanone (NaOC ₂ H ₅) Note: References 206-229 are fisted on p. 197.

TABLE VI-Continued

O ASSESSMENTONES	WITH P-raminotation
TABLE VI	ACTIVE METHYLENE COMPOUNDS WITH P-AMINOLOGY

Yield Refer-	enco 215	584	24		\$	
Yield	%	l	36		21	
	Cyelized Produet	ł	_		CH3	>
	Cyclising Agent —	kyla- tion failed 50 —	1		1	
	Yield %	kyla- tion failed 50	1		1	
ALKYLATIONS OF ACTIVE	Simple Alkylation Product	ſ	H,CC, CH,CH,CO,H	I		
ALKYI	Active Methylene Compound (Con- densiag Ageat)	Ω	Diethyl malonato (NaOH sus- pended in xyleno)	CH ₂ N(CH ₃) ₂ Ethyl acetoacetate (NaOC ₂ H ₅)	Ethyl methyl- acetoacetato (NaOC2Hs)	
	\$ 10 mm	g.Dimethylaminomethyl- 6-methyloyclohexanono	2.Dimethylaminomethyl-Diethyl malonako 6.phenyloyelobexanono (NaOH sus- pended in xyleno)	CH ₂ N(CH ₃) ₂	CH ₂ N(CH ₃) ₂ Ethyl methyl- acetoacetato (NaOC ₂ H ₃)	

Note: References 206-229 are listed on p. 197.

TABLE VII

Alkylations of Alkali Cyanides with β -Aminoketone Hydrochlorides

β-Aminoketone (as hydrochloride)	Product	Yield %	Refer ence
• • • •			
β-Dimethylaminopropiophenone	β-Benzoylpropionitrile	67	13
β-Dimethylamino-4-chloropropiophenone	β-(4-Chlorobenzoyl)propionitrile	32	13
β-Dimethylamino-4-hromopropiophenone	β-(4-Bromobenzoyl) propionitrile	63	13
β-Dimethylamino-3-nitropropiophenone	Resins		13
β-Dimethylamino-3-hydroxypropiophenone	$oldsymbol{eta}$ -(3-Hydroxyhenzoyl)propionitrile		13
β-Dimethylamino-4-hydroxypropiophenone	β -(4-Hydroxyhenzoyl) propionitrile	59	13
β-Dimethylamino-3-methoxypropiophenone	β-(3-Methoxyhenzoyl)propionitrile	73	13
β-Dimethylamino-4-methoxypropiophenone	β -(4-Methoxybenzoyl) propionitrile	71	13
β-Dimethylamino-3,4-dimethoxypropio- phenone	β-(3,4-Dimethoxybenzoyl)propio- nitrile	85	13
β-Dimethylamino-3,4,5-trimethoxypropio- phenone	β -(3,4,5-Trimethoxybenzoyl)- propionitrile	65	216
β-Dimethylamino-4-methylpropiophenone	β-(4-Methylbenzoyl)propionitrile	52	13
α-Dimethylaminomethylpropiophenone	Resin or oil	_	13
β-Dimethylaminopivalophenone	Isobutyrophenone	68	11
β-Dimethylaminoethyl α-naphthyl ketone	β-(1-Naphthoyl)propionitrile	43	13
β-Dimethylaminoethyl β-naphthyl ketone	β -(2-Naphthoyl)propionitrile	38	13
2-Dimethylaminomethylcyclohexanone	Resin or oil	_	13
β-Dimethylaminoethyl 2-furyl ketone	β-(2-Furoyl) propionitrile	57	13
β-Dimethylaminoethyl 2-thienyl ketone	β-(2-Thenoyl) propionitrile	67	13
β-Dimethylaminoethyl 2-henzofuranyl ketone	β -(2-Coumarilyl) propionitrile	21	13

Note: References 206-229 are listed on p. 197.

TABLE VIII

Carron Alkylations with Methiodides of β -Aminoketones

Physical Conference Action Metally less Compound Simple All-Jultion Product Simple All-Jultion			C	RGANIC	REAC	TIONS			
Simple Altylation Product 75	Reference 68	37	32, 32a	217		217a	25, 218	25	219
Simple Alkylation Product 776 Agent Rent 776 Agent Rent	 X2614	١	20	10	1	22	59	38	i
Simple Alkylation Product % % % % % % % % %		3-Methyl-4-carbethoxy-2-oyclobexen-	3-Methyl-8-isopropyl-2-cyclohexen- J-one (piperitone)	' ' '		\ //		10-Methyl-2-keto-∆¹. 9-octalin	1-Methyl-2,5-diketo-∆ ^{1,2} -octalin
Simple Alkylation Product freil) CHICOCHICO-CHGO-CHGIA CHICOCHICO-C-GAICH-CHGO-CHGIA 14ste	Cyclining Agent	NaOC ₂ H6	кон	1		1	$NaOC_2H_b$	1	7.7
Sention of the senting of the sentin	Yield % 17	27	1			1	1	1	1
Freil)	Simple Alkylation Product	CH,COCH(CO,C,U,)CH,CH2COCH1	t	1		i			O
A twinetrone as Methicular E-Merphalanone E-Merphalanone E-Merphalanone E-Merphalanone E-Merphalanone E-Merphalanone E-Merphanone E-Merphalanone E-Merphalan	Active Methylene Compound (Condensing Arent)			(NaoCell) (NaOCell) (NaOCell)		CH(CO,N)CH,CO,R CH,CH,CO,R (NaOCH,) (R = CH, C,H)	t-Methylcyclopentanono (NaNH3)	-Methyleycloberanone	.3-Cyclobernoedions (1)
	for Aminol ctorio	J. Mer Coning	2-Dietayismino- 3-butanone 1-Morrholmo-	Flutanons I. Dichy lamino- I-futanone		1-Diethylamino- 3-bulanono	I-Diethylamino- ? 3-butanono	1.Diethylamioo-	

f-Dictiyjamino- 2-Methyl-1,3-cyclohexanedione 3-butanone (NaOCH3)	Í	1	ı	H_3G CH_3	20	220
2-Carbethoxycyclobexanone (NaOC ₂ H ₅)	Í	f	ţ	H ₂ CH ₂ CO ₂ H I0-Carbethoxy-2-keto-∆ ^{1, 9} -octalin	38	25
2 -Carbomethoxycycloheptagone (NaOCH $_3$)	(CH2)4 CO CH2	86	кон + н,о + сн,он	CH (CH ₂) ₅ —CH CH ₂	65	27
1-Djethylamino- 2-Carbomethoxyeyoloğetanone 3-butanone (NaOCH4)	COSCH3 (CH2)s COCH3 (CH2)s COCH3 CCH2 COCH3	48	HC! + CH ₃ CO ₂ H	CH ₂ CH CH CH (CH ₂) ₅ —CH CH ₂ CH CH ₂ CH CH ₂ CH CH ₂ CH	14	88
				(CH ₂) ₅ ¢0 ¢H ₂	32	
1-Diethylamino- 2-Carbomethoxysyclononanone 3-butanone (NaOCH3)	CH2)8 CO CH2 CH2)8 CO CH2	79	HCl + CH3CO2H	CH2)s CO CH2 CH2)s CO CH2 CH2)s CO CH2	92	28
1-Dicthylamino- 2-Carbomethoxycyclodecanone 3-butanonc (NaOCH3)	CO ₂ CH ₃ CH ₂ COCH ₃ (CH ₂) ₇ CO CH ₂ (CH ₂) ₇ COCH ₃	18	HCI + CH3CO2H	(CH ₂) ₇ CO CH ₂	02	28
1.Diethylamino- 2.Carbomethoxyaxickridecanone 3.bulanone (NaOCH3)	CO ₂ CH ₃ (CH ₂) ₁₀ CO CH ₃ (CH ₂) ₁₀ CO CH ₂	1	HCI + CH ₃ CO ₂ H	(CH2)10 CO CH2	1	29
Note: References 206-229 are listed on p. 197.	солсня			24-0112		

TABLE VIII-Continued

Refer- ence	ā	21	221	221	221	221		221		221	
Yield]		81	11	48 ‡	Quant. (crude)	12-20 ‡		47 (72 ‡)		68 (82‡)	
KETONES Cooling Product	(CH2)12 CO CH2	(CH2)12 CO CH2	CH_{2} G-Cyclohexyl-10-carbomethoxy-2-keto- Δ^{1} 9-octalin	6-Cyclohexyl-2-keto-41, 9-octalin	6-Cyclohexyl-2-kcto-∆ ^{1,9} -octalia	6-Cyclohexyl-2-keto-A ^{1, 9} -oetalin		6-Cyclohexyl-2-keto-A ^{1, 9} -octalin		6-Cyclohexyl-2-keto-∆¹, 9-ootalin	
OF B-AMINO Cyclizing	Ageot HCl+ CH3CO2H	кон + H2O + СН3ОН	NaOCH3	7 802	HCI+	HOM +		HCI + CH,CO2H		кон + н20	
IODIDES	788		94			i		76		21	
CARBON ALKYLATIONS WITH METHIODIDES OF \(\beta\)-AMINOKETONES OF \(\beta\)-AMINOKETONES	Simple Alkylation Product ———————————————————————————————————	coch;	со,сн,	H ₂		CH	nooriosi 10N	СНО	$CH_2 \longrightarrow C_6H_{11}$	— — Сн. Сн.	CH2 OOH11
Сав	Activo Methylene Compound (Condensing Agent) 2-Carbomethoxycyclopentadecaoono (NaOCH3)			CO2CH3	Coduin	(NaOCH ₃)	COCO ₂ CH ₃		$HOCH = C_6H_{11}$	(NaOCH ₃)	
	p-Aminoketone as Methiodido 1-Diethylamino- 3-butanono			1-Dimethylam- ino-3-buta-	none		1-Dimethylam- ino-3-buta- 000e		1-Dimethylam- ino-3-buta- nooe		

		CARBON	ALKYLATIONS	WITH	AMINES	185
301	302	302	304	ន	222	123
83-90 73 over- all	l	ī	1	61	E3	σ
ПО200	HO ₂ C	HO I	H3CO2C			H ₃ C
1		1	١	1	1	1
88-18		91-100	1	ı	1	ī
CHO CHO_CH3_CH3_COCH3	HO ₂ CCCH ₃	CHO CHO CHO CHO CHO CH 2 CO CH 3	H ₃ CO ₂ C CH ₃ COCH ₃ CH ₃ CO ₂ CO ₂ COCH ₃ CH ₃ CO ₂ COCH ₃ C	i	I	i
Снон		нон)=0 =CHOH	(stereoisomer of above) (NaOH) 1-Diethylamino- frons-8-Decalone 3-butanone	1-Diethylamino- 2-Keto-A ^{9, 10} -octalin 3-butanone	1-Diethylamino- 3-butanone H ₃ C O= (NaMH ₂) Note: Reference 206-229 are listed on p. 197.

TABLE VIII-Continued

Methiodides of β -Aminoketones

Refer-	218	. 555	224	224	6 2
Yield	2 23	Ħ	28	16	1
THE TAX TO	Oyelized Fronce CH ₃		H,C OCH,	H ₃ C CO ₂ C ₂ H ₆	H ₃ C
Gvelizing	Agent	КОН	1	I	ı
IDES OF	268	1	1	1	1
CARBON ALEXTRATIONS WITH METHIODIDES OF B-AMINORETORIES.	Simple Alkylation Product	E C	і Н О	I	Ĭ
Сливо	7	(NaMHs) (Namh	11,10 00H3		(MaNH2) (MaNH2) (MaNH2)
	g.Aminoketono na Methiodido I.Diethylamino- 3-butanono	1-Diethylamioo- 3-butanono	1.Diethylamino- 3.hutanone	1-Diethylamino- 3-butanono	1-Diettylamino- 3-butanono

	CARBON	T ALKYL	ATIONS WI	TH AMINES		187
302	304	304	30a 31	25	ક્ષ	
80-88 (72 over- all)	æ	20		×	12	
По20	Ho;c	0-00°C	но-с	O H ₃ C OCH ₃	H,C OCH,	
1	1	1	1	1	1	
CHO CH2 CH2 COCH3	HO2C CH2CH2COCH3 51 CH2 CH2COCH3	H02C	H ₃ CO ₂ C ¹	1	1	
3-butkanone HO ₂ C	(NaOH) Diethylamino- 3-butanone	HO ₂ C (NaOH) (NaOH) 2-brithylamino- 2-brithylamino-	1-Diethylamino-		Diethylamino-	Note: References 206-229 are listed on p. 197.

TABLE VIII-Continued

Carron Alexidations with Methiodides of eta-Aminoketones

Refer-	220	80	ဗ	80 80	ii
Yield	43-48	ı	t	t	1
	CH ₃ O CH ₃ O CH ₃	I	CII, OCII,	CH, OCH,	CH,
7	Agent Agent	1	ì	1	-
ט פשעונעט	Xeld	ŧ	i	t	5 .
CARBON ALKYLATIONS WITH IMETHIODIDES OF P. 12.12.	Simple Alkylation Product	CH ₃ COCH ₂ CH ₃ COCH ₃)	t	CH,COCH,CH, CO,CH,
CARB	Active Methylene Compound (Condensing Agent) CH ₂ OCH ₃	$\begin{array}{c} \operatorname{dch}_{1} \\ \operatorname{ch}_{3} \operatorname{c} \\ \operatorname{ch}_{3} \operatorname{c} \\ \operatorname{ch}_{5} \operatorname{c}_{2} \end{array}$	(large eloess NaNH2)	CH ₃ OCH ₃	(NaNH2) Methyl fluoreno-9-carboxylate (NaOCH3)
	B-Aminotetone as Methiodide 1-Diethylamino- 3-butanone	1-Diethylamino- 3-butanone	e.	1-Diethylamino- 3-butanone	1-Morpholino- N 3-butanone

1-Diethylamino-3-pentanone

	CARBO:	N ALKYLA	TIONS WIT	II MIIII
26	26	26	83	33
73	06	£	20	31
CO ₂ CH ₃			CH ₂	
92 NaOCH3	KOH + CH30H	HCI + CH,CO2H	29 NaOC2Hs	н.50.
CO ₂ CH ₃ CH ₂ CH ₂ COCH ₃			CH ₂ CH ₃	C_2H_5 C_{B_3}
CO2CH2	NaOCHs)		H ₃ C	(NaNH2)

1-Dimethylamino-3-butanone

TABLE VIII—Continued

Rofer-	cneo		218		32, 32a	32, 32a	32	324	324	27a	
Yield	% 1		1		83	12	1	45	45	15	
Toketones	Cyclized Product	OCH ₃	CH ₃	H ₃ C	3,4,6-Trimetlyyl-2-cyclohexen-1-one	3-Isopropyl-G-methyl-2-cyclohexon- I-ono (carvenone)	2-Phanyl-O-koto-1-cyclohozeneaeetio acid	3.4.Butyl-2-oyelohexen-1-one	3-Isobutyl-2-oyelohexen-1-one	HO OH	>
of β -Amin	l Cyolizing Agent NaOC ₂ Hs				ном	кои	1	кон	КОН	псі + сп,со ₂ н	
BEGIOC	Yield % 23				1	1	1	l	١	20	
Carbon Alkylations with Methiodides of eta -Aminoketones	Simple Alkylation Product ÇII.a	CH3CH2COCH2CH3	or CH ₃		cn,cn,cocu,cu,	1	1	l	ı	O CH2	(CH ₂) ₄ —C
CAR	Activo Methylens Comp (Condensing Agent)	E C	(NaNH2)		mit	Ethyl methylacetoacetate Ethyl methylacetoacetate		(NnOC2H5) Ethyl nectoacetato (NnOC2H5)	Talend and anadatala	Etnyi mecuatette (NaOC2Hs) 2-Carbethoxyeyelohexanono (NaOCH3)	
	A-Aminoketono as Methiodido	1-Diethylamino- 3-pentanona			•	1-Morpholino- 2-methyl- 3-butanone	4-methyl- 3-pentanone 1-Moreholino-	3-pentanone 1-Morpholino-	3-pentanono	I-Morpholino- 5-methyl- 3-hexanone 1,1-Bis(diethyl- nmino-	methyl)- acctono

	CAR	BON ALKY	LATION	s with	AMINES	
272 a	27.0	27a	27.0		27a	27.0
61	1	35	15	ā	37	S
69 HCl+Co ₂ H CH ₃ CO ₂ H	1 1	63 HG+CO ₂ H (CH ₂), HO CH ₃ CH ₃	CHCH2) CH3	68 HCl.+ CH2.0 ₀ H	CI(CH ₂) CH ₃	CH3CO ₂ H (CH ₂) ₁₀ I
O CH2	CO2CH3	CO ₂ C ₃ H ₈ CO ₂ C ₃ H ₈ CH ₂ CCOCH ₃ CH ₂ CCOCH ₃	CO ₂ CH ₃	CH2CCCH3	CO ₂ OH ₂	O CH2
1,1-Bis (diethyl- 2-Carbethoxycyclohcptanone amino- methyl)-		1,1-Bis(diethyl- 2-Carbomethoxycyclooctanone amino-methyl)- actone		1,1-Bis(diethyl- 2-Carhomethoxycyclonounoue amino- methyl)- acetone		1,1-Bis(diethyl- 2-Carbomethorycyclotridecanone amino- methyl)- acetone
1,1-Bis(diethyl- amino- methyl)-	acetone 1,1-Bis(diethyl- amino- nethyl)- acetone	1, I-Bis(diethy amino- methyl)- acetone		1,1-Bis(dieth amino- methyl)- acetone		1,1-Bis(die amino- methyl). acetone

Note: References 206-229 are listed on p. 197.

TABLE VIII—Continued

	β -Aminoketones
	OF
יייייט ייייי	ALEYLATIONS WITH METHIODIDES OF B-AMINOKETO
	WITH
47	ATENTATIONS
	7,000

Reference 27	26	26	37	37	37	37
Yield % 80	43	70-83	1	1	١	13
Cyclized Product $(CH_2)_{12} \to O$ CH_3	H ₃ C H ₁₀ C	HG + CH ₃ CO ₂ H H ₃ C CH ₃ CO ₂ CH ₃ CO ₂		3-(2'-Dimethylamino)ethyl-6-car- bethoxy-2-cyclobexen-1-ons	ı	CH2N(CH3)2
Cycling Agent HCI+ CH3CO2H	кон , Сн ₃ он	HCl + CH ₃ CO ₂ H then (CH ₃ CO) ₂ C	1	H ₂ SO4	1	BCI
Yield % 58	22		22	21	1	ro.
CARBON ALKYLATIONS WITH IMETRICIAL CONTROL OF CHAPTER O	CH_3O_2C CH_3O_2C $CH_2=C$ $CH_2=C$ $O=C$	CH ₃	COCH ₂ CH ₂ N(CH ₃) ₂	ĊĦ¿CĦ¿CH(CO¿C¿Ħゟ)₂ CĦţCOCHCO¿C¿Ħゟ	CH2CH2COCH2CH2N(CH3)2 CH3CCH(CO2C2H5)CH2CH3]2CO	OO-00 (CH ₃) ₂ N(CH ₂) ₂ CO(CH ₃) ₂ COO.H-
Active Methylene Compo (Condemning Agent) 2-Carbomethoxycycloputad (NaOCH3)	сы, о, с	(NaOCH ₃)	Diethyl malonate	Ethyl acetoacetate	(ACCARS) Ethyl acetoscetate	2-Carbethoryeydopentanone (KOC2H\$)
A-Aminoketone as Methiodide 1,1-Bis(diethylanino-methyl)- soctone	1,1-Bis(di- methylamino- methyl)- acetono		1-Methyl-			

	CARBO	N AL	KYLATI	ONS WI	IH AMIM	BO
37	32,32a	32	227	227	Ħ	215
1	3	1	1	1	1	1
1	3-Phenyl-2-cyclohexen-1-one	2.Phenyl-6-keto-1-cyclohexeneacetic acid		C ₆ H ₆	1	I
i	H	1	6 <i>0</i> 1	1	1	1
72	— кон	1	83	88	20	09
·						
CH_{2} CH_{2} CH_{2} CH_{3} CH_{2} CH_{3} CH_{4}	CH ₂ N CH ₃	1		Cohicochichi Colchi	C ₆ H ₆ COCH ₂ CH ₂ CO ₂ CH ₃ β-Dimethylaminopivalophenone	CH2CH(CO2C2H4)2 =0 CH2CH2CO2C2H5
1-Ethyl- 4-piperidooe 2-Carbethoxyoyolohoxanono 4-piperidooe		g-Morpholino- Ethyl acetoacetate propio- phenono	β-Morpholino- COCH ₂ CO ₂ C2H b propio- CH ₂ CH ₂ CO ₂ C2H, phenone CH ₂ CH ₂ CO ₂ C1, β-Diethylamino- Methyl fluorene-3-carbax/late propio-	phenone (metho- suliate) «Mornholino- Methyl fluorenc-9-carboxylate	propolo- propolo- propolo- prenone (methosulfate) β-Dimethyl- Sodium cyanide	aminopivalo- phenone 2-Dimethylam- Diethyl malonato inomethyl- cyclohexanone

Note: References Wo-229 are listed on p. 197.

TABLE VIII—Continued

	th Methiodides of eta -Aminoketones
3	OF
THE COMME	HIODIDES
111	MET
	WITH
47	CARBON ALKYLATIONS WITH M
	CARBON

Refer- ence 25 215			215		23	224	221	224
Yield % P0		1	ı		8	27	23	13
Cyclized Product 2-Keto-A ^{4, 0} -octalin		I	1		2-Keto-8-methyl-A ^{1, 9} -octalin	2-Keto-6-methoxy- $\Delta^{1,9}$ -octalin	2-Keto-1-methyl-6-methoxy-A1-9-octa-	2-Keto-1-methyl-6-carbomethoxy- At 9-octalin
Cyclizing Agent		1	1		1	1	ı	ì
Yield %	2	28	42		ı	1	ı	1
Simple Alkylation Product	H,CC CH2CH(CO2C2H6)2 =0	H3CCH2CH2CO3C3H5	CH ₂ CH(CO ₂ C ₂ H ₅) ₂ CH ₃	CH2CH2CO2C3U5 =0 CH3	ı	1	ı	I
Active Methylono Compound (Condensing Agent) Ethyl acetosciate (NaOC2H5)	Diethyl malonate (NaOC2Hs)		Diethyl malonate (NaOC ₂ H ₆)		Ethyl acetoacetate	Ethyl acetoacetate	Ethyl propionylacetate	(NaOC2H5) Ethyl propionylacetate (NaOC2H6)
p-Aminoketone as Methiodide 2-Diethylaminomethylgyclo- bexanone	2-Dimethylaminomethyl-4- methylcyclohexanone		2-Dimethylamiromethyl-G- methylcyclobexanone		2-Diethylaminomethyl-6-methyl- Ethyl acetoacetate			oxycyclonexanone 2-Diethylaminomethyl-4-car- bomethoxycyclobexanone

١

Ethyl acetoacetate (NaOC2Hs)

		0111	
32a	32, 228	32 32b 70	229
11	20	1 82 94	ध
4,4-Dicarbethoxy-1-trans-decalone	H ₃ C CH(CH ₃) ₂	2.Keto-5-methyl-8-isopropyl- 2.Keto-5-methyl-8-isopropyl-A ^{1,9} -octa- lin C ₂ H ₅ O ₂ C	
1	t	кон	1
1	1	1 1 1	1

١

Ethyl acetoacetate (NaOC2Hs) Ethyl acetoacetate (NaOC2Hs)

2.Piperidinomethyl-3-methyl-6-isopropylcycloheranone 2.Morpholinomethyl-3-methyl-6-isopropylcycloheranone

Ethyl acetoacetate

١

CH(CH₃)₂ Ethyl methylacetoacetate (NaOC₂H₆)

١

Diethyl malonate (NaOC2Hs) ١

Note: References 206–229 are listed on p. 197.

• The simple alkylation product was not isolated.

† The simple alkylation product was opticated as the methyl or isopropyl ether.

† The product was isolated as the semicarbaxone.

† The product was optized as the semicarbaxone.

§ The material was optized after decarbaxilation and reduction to the alcohol.

§ The is the combined yield of the two products.

TABLE IX

AN ALKYLATION OF INDOLE WITH DIETHYL PIPERIDINOMETHYLFORMAMIDO-MALONATE 69

		Yield of	Indole
		Alkylated	Mannich
		Product (A)	Base (B)
Solvent	Catalyst	%	%
Mesitylene	NaOH	10	0
Xylene	$N_{8}OH$	76	0
Xylene	None	0	70
Toluene	NaOH	21	0
Toluene	None	16	22
Benzene	$N_{8}OH$	0	13
Benzene	None	0	0

TABLE X

CARBON ALKYLATIONS WITH MANNICH BASES DERIVED FROM NITRO COMPOUNDS 210

	00			
Mannich Base or Quaternary Salt	Compound Alkylated	Solvent; Catalyst	Product	Yield % 34
1-Dimethylamino-2-nitro- hutane	1-Nitropropane	None; NaOH	3,5-Dinitroheptane	18
1-Diethylamino-2-nitro- hutane	1-Nitropropane	None; NaOH	3,5-Dinitroheptane	55
1-Dimethylamino-2-nitro- hutane	2-Nitropropane	None; NaOH	2-Methyl-2,4-dinitro- hexane	23
1-Diethylamino-2-nitro- hutane	Methyl cyanoacetate	Xylene; none	Methyl 2-cyano-4-nitro- hexanoate	16
1-Diethylamino-2-nitro- hutane	Ethyl cyanoscetate	Xylene; none	Ethyl 2-cyano-4-nitro- hexanoate	10
1-Dimethylamino-2-methyl- 2-nitropropane	2-Nitropropane	None; NaOH	Alkylation failed	
1-Dimethylamino-2-methyl- 2-nitropropane methiodide	Ethyl acetamido- cvanoacetate	_	Alkylation failed	
1-Dimethylamino-2-methyl- 2-nitropropane methiodide	α-Naphthol	_	Alkylation failed	

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CHAPTER 4

THE VON BRAUN CYANOGEN BROMIDE REACTION

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^{*} Present address: Naugatuck Chemical Company.

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INTRODUCTION

The reaction of a tertiary amine with cyanogen bromide was first described in 1900 by Julius von Braun,¹ who subsequently elaborated the reaction to such an extent that it rightfully bears his name. The reaction apparently was discovered independently by Scholl and Nörr,² whose paper was submitted for publication five weeks after the submission of von Braun's first paper.

Generally, a tertiary amine reacts with cyanogen bromide to yield an alkyl bromide and a disubstituted cyanamide. The direct conversion of

$$R''$$
 R'
 R'
 $N + BrCN \rightarrow RBr + NCN$
 R'

secondary amines to disubstituted cyanamides with cyanogen bromide proceeds in low yield because some of the amine is converted to its hydrobromide. Furthermore, the amine hydrobromide frequently reacts with the cyanamide formed to give a guanidine as the principal product.³ Preliminary conversion of the secondary amine to a tertiary amine by reaction with formaldehyde, followed by cleavage of the product with cyanogen bromide, affords a better yield of the disubstituted cyanamide.

An acyclic amine yields an alkyl bromide and a disubstituted cyanamide as discrete products. The bromide and cyanamide obtained from the cleavage of a monocyclic amine, such as an N-substituted pyrrolidine,

¹ von Braun, Ber., 33, 1438 (1900).

² Scholl and Nörr, Ber., 33, 1550 (1900).

³ von Braun, Ber., 42, 2035 (1909).

may be discrete compounds, or they may constitute portions of the same molecule. The product from a bicyclic amine necessarily contains

$$\begin{array}{c|c}
 & \longrightarrow & \longrightarrow & + RBr \\
 & & \longrightarrow & N \\
 & & & CN \\
 & & & RN(CN)CH_2CH_2CH_2CH_2Br
\end{array}$$

the bromine and the cyanamide group in the same molecule. Nitrogen heterocycles such as pyridine add a mole of cyanogen bromide at the carbon-nitrogen double bond.

$$\begin{array}{c|c}
& \xrightarrow{BrCN} & & H \\
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An elimination reaction resulting in the formation of an olefin can occur.4,5 The presence of a secondary or tertiary alkyl group in the

$$\begin{array}{c} R' \\ C(CH_3)NR''R''' \xrightarrow{BrCN} \\ R' \\ C=CH_2 + NCN + C(CH_3)NR''R''' \cdot HBr \\ R \\ \end{array}$$

amine is conducive to olefin formation. When the reaction takes this course, a considerable quantity of the amine is converted to the hydrobromide and is thereby prevented from reacting with the cyanogen bromide.

Von Braun 4 early in his work noted differences in the vigor of the reaction of various amines with cyanogen bromide. Simple aliphatic amines react so vigorously that dilution with an inert solvent is required to keep the reaction under control. Derivatives of aniline react less readily; N-alkyldiphenylamines require relatively strenuous conditions for cleavage and give poor yields of products. As the nucleophilic character (basicity) of the nitrogen atom is reduced, its tendency to react with cyanogen bromide is lowered; e.g., N-substituted amides do

⁴ von Braun, Ber., 33, 2728 (1900).

⁵ Elderfield and Hageman, J. Org. Chem., 14, 605 (1949).

not react with cyanogen bromide. The nucleophilic strength of cyanamides is sufficiently low to prohibit reaction with cyanogen bromide.6 Consequently, when an amine is cleaved by cyanogen bromide, there is no danger of any subsequent reaction between the cyanamides formed and excess cyanogen bromide.

The most thoroughly investigated aspect of the von Braun cyanogen bromide reaction is its use to establish the relative lability of various carbon-nitrogen bonds in tertiary amines. For this purpose it is necessary to determine which of the three substituents is displaced as an alkyl bromide or olefin. Correlation of the large amount of data on the relative ease of removal of different groups enables one to predict approximately how a particular amine will be cleaved (see p. 231 and Table I). Depending upon the structure of the amine, the cleavage may proceed entirely in one direction, or it may give a mixture of all possible alkyl bromides and disubstituted cyanamides.

A serious interfering side reaction involves reaction of the amine with the alkyl bromide produced by the cleavage to form a quaternary ammonium bromide. This side reaction, which is particularly serious when

$$\begin{array}{c}
R'' \\
2 R' \longrightarrow N + BrCN \longrightarrow \begin{bmatrix} R' \\ N(R'')_2 \end{bmatrix} Br * + NCN \\
R
\end{array}$$

highly reactive bromides are involved, is minimized by making certain that the amine is continually in the presence of excess cyanogen bromide during the reaction.

A survey of the literature discloses surprisingly few cases in which the von Braun cyanogen bromide reaction has been employed for synthetic purposes. It has been applied mainly as a method of degradation in the structural analysis of alkaloids.

Many cleavages, however, when run under the proper experimental conditions, proceed smoothly, and the products are obtained in excellent It appears that the reaction could be applied more widely in synthetic organic chemistry than it has been (see p. 224). Unfortunately, much of the experimental work reported lacks details, particularly with regard to yields, and this may have prevented the reaction from attaining wider synthetic use. Many of the reactions reported to give a mixture of products in low yield could certainly be improved by the proper choice of experimental conditions.

von Braun, Ber., 36, 2286 (1905).
 For convenience, ionic charges will not be shown when it is obvious that the substance represented is a simple quaternary salt.

The material in this chapter is limited to a discussion of the reaction of tertiary * amines with cyanogen bromide. Reactions of cyanogen bromide with other compounds are mentioned only when they add to this general discussion. The effect of the structure of the amine on the direction of cleavage by cyanogen bromide is emphasized.

MECHANISM

Von Braun's observation of the formation of an initial, transient precipitate 1,7 when an amine is mixed with cyanogen bromide led him to propose the preliminary formation of an unstable complex involving quaternary nitrogen. This intermediate is stable only at low temperatures and has never been isolated for characterization.

A brief consideration of the structure and chemical behavior of cyanogen bromide is helpful in understanding its reaction with amines. On the basis of X-ray diffraction studies 8 and Raman spectral data,9 cyanogen bromide has the structure Br—C≡N rather than Br—N≡C⁻. In the cyanogen halide series cyanogen chloride nearly always reacts with displacement of the chlorine as chloride ion, whereas in cyanogen iodide the presence of positive iodine is indicated.10 Cyanogen bromide occupies an intermediate position with respect to the polarity of the carbon-halogen bond. The brominating action of cyanogen bromide " and its reaction with Grignard reagents 12 suggest the presence of a positive bromine atom. However, in the greater number of reactions of cyanogen bromide the bromine is displaced as bromide ion. Reaction with aqueous alkali forms bromide and cyanate ions.10 Reaction with aqueous solutions of primary, secondary, or tertiary amines yields bromide ion quantitatively.13 The electrolysis of cyanogen bromide in a variety of organic solvents results in migration of bromine to the anode as bromide ion.14

The initial reaction of cyanogen bromide with an amine involves a displacement of the bromine as bromide ion with the formation of an

⁷ von Braun, Ber., 40, 3914 (1907).

10 Kleinberg, J. Chem. Education, 23, 559 (1946).

¹² Grignard, Bellet, and Courtot, Ann. chim., [9] 4, 28 (1915).

^{*} Throughout the remainder of this chapter the word "amine" is used to designate a tertiary amine unless otherwise indicated.

⁸ Pauling and Hendricks, J. Am. Chem. Soc., 48, 641 (1926).

⁹ West and Farnsworth, J. Chem. Phys., 1, 402 (1933).

¹¹ Migrdichian, The Chemistry of Organic Cyanogen Compounds, p. 115, Reinhold Publishing Corp., New York, 1947.

¹³ Griffith, Jobin, and McKeown, Trans. Faraday Soc., 34, 316 (1938). ¹⁴ Clark and Streight, Trans. Roy. Soc. Can., [3] 22, III, 323 (1928) [C. A., 23, 1824 (1929)].

ionic addition compound in which the nitrogen atom is quaternized. As the terminating step, a nucleophilic displacement by bromide ion removes one of the substituents as an alkyl bromide. Von Braun 4

$$\begin{bmatrix} R'' \\ R' - NCN \end{bmatrix}^{+} + Br^{-} \rightarrow R'' NCN + RBr$$

defined the vigor of the reaction as the ease of formation of the quaternary compound. Reduction of the nucleophilic strength of the amine decreases the readiness with which the addition compound is formed. This mechanism is compatible with the known ability of quaternary ammonium salts to function as alkylating agents.15 The elimination reaction that has been observed 5 when an amine containing a secondary or a tertiary alkyl group is treated with cyanogen bromide can be interpreted in a manner consistent with this mechanism.

No kinetic studies of the von Braun cyanogen bromide reaction have been reported that shed any light on the mechanism under the conditions normally employed. In fact the only recorded kinetic study of the reaction of cyanogen bromide with amines deals with a measurement of the rate of formation of bromide ion in aqueous solution.¹³ Although second-order kinetics were observed in aqueous solution, the course of the reaction in this instance is admittedly not identical with that in a non-polar solvent.

Evidence supporting a mechanism involving a second-order displacement by bromide ion is afforded by the observation that those alkyl groups whose halides are known from other studies to react readily in displacement reactions are also most readily cleaved from amines as alkyl bromides.16

In this formulation, the von Braun reaction is akin to other reactions of tertiary amines characterized by conversion of the nitrogen to the quaternary state, followed by dealkylation. Some examples follow.

(a) Acetyl bromide reacts 17 with dimethylaniline in much the same manner as does cyanogen bromide. The formation of the disubstituted

manner as does cyanogen by
$$2C_6H_6N(CH_3)_2 + CH_3COBr \rightarrow [C_6H_5N(CH_3)_3]Br + CH_3CON(CH_3)C_6H_6$$

$$2C_6H_6N(CH_3)_2 + CH_3COBr \rightarrow [C_6H_5N(CH_3)_3]Br + CH_3CON(CH_3)C_6H_6$$

¹⁵ Snyder, Smith, and Stewart, J. Am. Chem. Soc., 66, 200 (1944); Snyder and Speck, Snyder, Smith, and Stewart, J. Am. Soc. chim. France, 39, 305 (1926); 45, 109 (1929). ibid., 61, 688, 2895 (1939); Rodinov, Bull. soc. chim. France, 39, 305 (1926); 45, 109 (1929). See also Chapter 3.

¹⁵ von Braun and Engel, Ann., 436, 299 (1924).

¹⁷ Stadel, Ber., 19, 1947 (1886).

acetamide is analogous to the formation of cyanamides by cyanogen bromide; both reactions form methyl bromide which may appear, as above, in a quaternary salt of the amine. Acyl chlorides undergo this reaction far less readily than acyl bromides.

(b) The dealkylation of an amine by a carboxylic acid proceeds much less readily than by an acid halide or anhydride. 18 Heating dimethylaniline to 210–220° with β -phenylpropionic acid gives a 15% yield of the

disubstituted amide.19

$${\rm C_5H_5N(CH_3)_2} + 2{\rm C_6H_6CH_2CH_2CO_2H} \xrightarrow{\rm Heat}$$

$$C_6H_6CH_2CH_2CON(CH_3)C_6H_6 + C_6H_6CH_2CH_2CO_2CH_3$$

(c) Demethylation of dimethylaniline is effected by heating with n-amyl bromide 20 or phenacyl bromide.21 These two reactions merely

$$C_6H_6N(CH_3)_2 + n \cdot C_5H_{11}Br \xrightarrow{150-160^{\circ}} C_6H_5N(CH_3)C_6H_{11} \cdot n + CH_3Br$$
 $C_6H_6N(CH_3)_2 + C_6H_6COCH_2Br \xrightarrow{70^{\circ}} C_6H_6N(CH_3)CH_2COC_6H_5 + CH_3Br$
convert one tertiary amine to another; in this respect they differ from

the other examples.

Cyanogen bromide reacts with thio ethers and with tertiary phosphines, arsines, and stibines in much the same way as with amines. Thio ethers undergo cleavage with the formation of an alkyl bromide and a thiocyanate, 22, 23, 24 but no analogous reaction has been observed with

$$RSR' \xrightarrow{BrCN} RSCN + R'Br$$

ethers. With thio ethers the relative ease of removal of various alkyl

groups parallels closely that observed with amines.

In contrast to triphenylamine, triphenylphosphine forms an addition compound with cyanogen bromide, but no cleavage to bromobenzene takes place. Phosphines appear to be attacked more readily by cyanogen bromide than are amines.25

$$(CH_3)_2N \xrightarrow{P(C_5H_5)_2} \xrightarrow{BrCN} Oil \xrightarrow{H_2O} (CH_3)_2N \xrightarrow{P(C_6H_5)_2 \cdot H_2O}$$

¹⁸ Tiffeneau and Fuhrer, Bull. soc. chim. France, [4] 15, 163 (1914).

¹⁹ von Braun and Weissbach, Ber., 63, 489 (1930). ²⁰ Claus and Rautenberg, Ber., 14, 622 (1881).

²¹ Stadel and Siepermann, Ber., 14, 984 (1881).

²² von Braun and Engelbertz, Ber., 56, 1573 (1923).

²³ von Braun, May, and Michaelis, Ann., 490, 189 (1931).

²⁴ von Braun and Friedsam, Ber., 63, 2407 (1930). 25 Steinkopf and Buckheim, Ber., 54, 1024 (1921).

Tertiary arsines react with cyanogen bromide 26,27,28 to form addition products that are considerably more stable than those from amines; for example, ethyldiphenylarsine yields an addition complex that can be isolated and undergoes cleavage only when heated.29 Tertiary stibines 30 react with cyanogen bromide in a similar manner.

$$(C_6H_5)_2A_5C_2H_5 \xrightarrow{BrCN} (C_6H_5)_2A_5(CN)C_2H_5]Br \xrightarrow{140^{\circ}} (C_6H_5)_2A_5CN + C_2H_5Br$$

SCOPE AND LIMITATIONS

Acyclic * Amines

The cleavage of an unsymmetrically substituted amine of low molecular weight occurs predominantly in the direction involving displacement of the smallest group.1 Upon ascending the normal aliphatic series,

$$(n-C_3H_7)_2NCH_3 \xrightarrow{BrCN} (n-C_3H_7)_2NCN + CH_3Br$$

 $(n-C_3H_7)_2NC_2H_5 \xrightarrow{BrCN} (n-C_3H_7)_2NCN + C_2H_5Br$

the ease of removal of the alkyl group decreases, the difference between adjacent homologs being greater between the lower members of the series. Above n-hexyl there is no detectable difference in the ease of cleavage of consecutive members.31 Other structural features, such as branching of the chain and the presence of β,γ -unsaturation, are far more significant than the size of the group. Cleavage of an aromatic amine to give an aryl bromide has never been observed. A rule that is helpful, though not inviolable, for predicting which alkyl group will be removed from the amine can be derived from a comparison of the relative reactivities of the corresponding alkyl bromides. Generally those groups, such as allyl and benzyl, whose halides are known to be highly reactive in displacement reactions 16,32 are cleaved more readily than less reactive groups. However, when a substituent is cleaved with the formation of an olefin, this rule is not applicable.

²⁶ Steinkopf and Wolfram, Ber., 54, 848 (1921).

²⁷ Steinkopf and Schwen, Ber., 54, 2791 (1921).

²³ Steinkopf and Müller, Ber., 54, 841 (1921).

²² Steinkopf, Donat, and Jager, Ber., 55, 2597 (1922).

³⁰ Morgan and Yarsley, Proc. Roy. Soc. London, Series A, 110, 534 (1926).

^{*} Morgan and Yarsiey, 1766. 268. Solden that the nitrogen atom of the amine * The term "acyclic" is employed here to define that cyclic substituents are excluded is not part of a ring. It is not used in the strict sense that cyclic substituents are excluded. 31 von Braun and co-workers, Ann., 507, 1 (1933).

York, 1940,

Since a phenyl group is not removed from an amine by cyanogen bromide, dialkylanilines containing different alkyl groups have been employed extensively for dealkylation studies. The *n*-propyl group is removed more readily than the isopropyl group when *n*-propylisopropylaniline is allowed to react with a molar equivalent of cyanogen bromide

$$C_6H_5N(C_3H_7-n)C_3H_7-i \xrightarrow{BrCN} C_6H_5N(CN)C_3H_7-i + n-C_3H_7Br$$

at the temperature of the steam bath. The tendency of the isopropyl group to undergo removal by an elimination reaction has been obscrved in the reaction of diisopropylaniline with cyanogen bromide. In this reaction an appreciable quantity of diisopropylaniline hydrobromide is formed. Since isopropyl bromide did not react with diisopropylaniline under comparable conditions, it can be concluded that the isopropyl group is removed directly from the quaternary addition compound as propylene.

The greater lability of the n-butyl group compared with the isobutyl group has been shown by the cleavage of n-butylisobutylaniline.³³ Very

$$C_6H_5N(C_4H_9\text{-}n)C_4H_9\text{-}i\xrightarrow{BrCN}C_6H_5N(CN)C_4H_9\text{-}i+n\text{-}C_4H_9Br$$

little cleavage to give isobutyl bromide was observed. More remote branching of the chain, as in the isoamyl group and higher homologs, is much less influential.

 β,γ -Unsaturation. The labilizing effect of β,γ -unsaturation is demonstrated by the cleavage of methylallylaniline, and diethylcyclopentenylamine. No mention was made of the isolation of any cyclopentenyl

bromide in the latter reaction. It is not surprising that transfer of the unsaturation to a more remote position greatly reduces the lability, as has been shown by the cleavage of dimethyl-4-pentenylamine.³⁵ This

u von Braun and Murjahn, Ber., 59, 1202 (1926).

¹⁴ von Braun and Kühn, *Ber.*, **60**, 2551 (1927).

¹⁵ von Braun and Kohler, *Ber.*, **51**, 79 (1918).

reaction illustrates a common side reaction involving the formation of a quaternary ammonium bromide by the reaction of the liberated alkyl bromide with the amine. A determination of the structure of the quaternary bromide reveals the direction of cleavage of the amine.

Though the benzyl group ³⁶ is more susceptible to cleavage from an amine by cyanogen bromide than the methyl group, a phenethyl group ³⁷

$$\begin{array}{ccc} \mathrm{C_6H_5CH_2N(CH_3)C_6H_5} & \xrightarrow{\mathrm{BrCN}} & \mathrm{C_6H_5N(CN)CH_3} + \mathrm{C_6H_6CH_2Br} \\ \mathrm{C_6H_6CH_2CH_2N(CH_3)C_6H_6} & \xrightarrow{\mathrm{BrCN}} & \mathrm{C_6H_5CH_2CH_2N(CN)C_6H_6} + \mathrm{CH_3Br} \end{array}$$

is more resistant to cleavage. When removed further than the β position, the phenyl group exerts no labilizing influence.

The removal of an allyl group in preference to a benzyl group is demonstrated by the cleavage of allyldibenzylamine and allylbenzylaniline.³⁶ In these reactions the products contained only traces of benzyl bromide.

An interesting labilizing effect is associated with the presence of a cyclopropyl group. The cyclopropylmethyl group ³⁸ is more readily removed than a methyl group. It is, however, less readily removed

$$\begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{CHCH}_2 \text{N(CH}_3) \text{C}_6 \text{H}_5 \end{array} \xrightarrow{\text{BrCN}} \text{C}_6 \text{H}_5 \text{N(CN)CH}_3 + \text{CH}_2 \\ \text{CH}_2 \\ \text{CHCH}_2 \text{Br} \end{array}$$

than a benzyl group.

Amines containing the more readily displaced substituents do not necessarily react more vigorously with cyanogen bromide. For instance, tribenzylamine does not react with cyanogen bromide at room temperature; heating to about 70° is required to effect an appreciable rate of reaction.¹

Substituted Allyl and Benzyl Groups. Extensive studies have been made of the effect of substituents on the ease of removal of allyl 16,29 and benzyl 16,23,24,31,39,40 groups. The introduction of a chlorine or bromine atom into the β or γ position of the allyl group increases the resistance to cleavage to the extent that these groups are less easily removed than a benzyl group. The difference between the effect of bromine and that

$$C_6H_5CH_2N(CH_3)CH_2CBr$$
= $CH_2 \xrightarrow{BrCN}$

 CH_2 = $CBrCH_2N(CN)CH_3 + C_6H_6CH_2Br$

³⁶ von Braun and Schwartz, Ber., 35, 1279 (1902).

m von Braun, Ber., 43, 3209 (1910).

³⁸ von Braun, Fussgänger, and Kühn, Ann., 445, 201 (1925).

³⁹ von Braun, Kühn, and Weismantel, Ann., 449, 249 (1926).

⁴⁰ von Braun, Michaelis, and Spanig, Ber., 70, 1241 (1937).

of chlorine on the lability of substituted allyl groups is too small to be detected by the method of product analysis employed. However, a halogen in the β position has been shown to produce greater resistance to cleavage of the group than one in the γ position.³⁹ An increase in the lability of the allyl group is caused by a phenyl group in the γ position.¹⁶

The presence of halogen in the ring of the benzyl group influences the lability of this group in a definite way. With the exception of substitution by fluorine, which appears to exert little influence, the halogen-substituted benzyl groups show greater resistance to cleavage than the unsubstituted benzyl group. The lability of the substituted benzyl group decreases in the order $Cl > Br > I.^{31}$ With reference to position,

 $p\text{-ClC}_6\text{H}_4\text{CH}_2\text{N}(\text{CH}_3)\text{CH}_2\text{C}_6\text{H}_4\text{Br-}p \xrightarrow{\text{BrCN}}$

p-ClC₆H₄CH₂Br + p-BrC₆H₄CH₂N(CN)CH₃

 $m\text{-BrC}_6\text{H}_4\text{CH}_2\text{N}(\text{CH}_3)\text{CH}_2\text{C}_6\text{H}_4\text{Br-o} \xrightarrow{\text{BrCN}}$

m-BrC₆H₄CH₂Br + o-BrC₆H₄CH₂N(CN)CH₃

the lability decreases in the order para > meta > ortho. Variation of the position exerts a more pronounced influence than variation of the halogen. This is shown by the cleavage of o-chlorobenzyl-m-iodobenzyl-methylamine.³¹ In the examples cited, the occurrence of cleavage almost

 $m\text{-}\mathrm{IC}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{N}(\mathrm{CH}_3)\mathrm{CH}_2\mathrm{C}_6\mathrm{H}_4\mathrm{Cl}\text{-}o \xrightarrow{\mathrm{BrCN}}$

m-IC₆H₄CH₂Br + o-ClC₆H₄CH₂N(CN)CH₃

exclusively in the directions indicated shows that the differences in the lability of these substituted benzyl groups are quite pronounced.

Other substituents, like the halogens, decrease the lability of the benzyl group most effectively when in the ortho position. Variation of the lability with change in position is not so marked with the nitro group as with the halogens.⁴⁰ Qualitative evaluation of the effect of different substituents in any particular position upon increasing the resistance to cleavage of the benzyl group gives the following decreasing order of effectiveness: $NO_2 > CN > I > Br > Cl > H$. The acetamino group ⁴⁰ has been shown to increase the resistance to cleavage of the benzyl group to a greater extent than chlorine but no data comparing it with bromine and iodine are available. The nitro and cyano groups

 $p\text{-}\mathrm{CH_3CONHC_6H_4CH_2N(CH_3)CH_2C_6H_4Cl-}p \xrightarrow{\mathrm{BrCN}}$

p-CH₃CONHC₆H₄CH₂N(CN)CH₃ + p-ClC₆H₄CH₂Br

increase the resistance to cleavage of a benzyl group to a greater extent than any of the halogens, even when the latter are in the ortho position.⁴⁰

$$p-O_2NC_6H_4CH_2N(CH_3)CH_2C_6H_4Cl-o \xrightarrow{BrCN}$$

p-O₂NC₆H₄CH₂N(CN)CH₃ + o-ClC₆H₄CH₂Br

 $p\text{-NCC}_6\text{H}_4\text{CH}_2\text{N(CH}_3)\text{CH}_2\text{C}_6\text{H}_4\text{I-o} \xrightarrow{\text{BrCN}}$

 $p ext{-} ext{NCC}_6 ext{H}_4 ext{CH}_2 ext{N(CN)CH}_3 + o ext{-} ext{IC}_6 ext{H}_4 ext{CH}_2 ext{Br}$

However, no case has been reported in which the lability of a benzyl group has been reduced by a substituent on the ring to the extent that its resistance to cleavage equals that of a methyl group.

Substituents that labilize the benzyl group, listed in the order of decreasing effectiveness, are as follows: methoxyl > phenyl, cyclohexyl > p-xenyl > ethyl > methyl > $H.^{23,31}$ In this series also, a substituent in the ortho position produces a less labile benzyl group than when it is in the meta or para position. Though a methyl group in the para position labilizes the benzyl group, a methyl group in the ortho position does not. However, the o-methylbenzyl group is more labile than the p-chloro-

$$0-\text{CH}_3\text{C}_6\text{H}_4\text{CH}_2\text{N}(\text{CH}_3)\text{CH}_2\text{C}_6\text{H}_6 \xrightarrow{\text{BrCN}}$$

 $C_6H_6CH_2Br + o\text{-}CH_3C_6H_4CH_2N(CN)CH_3$

benzyl group. 31 The p-methoxybenzyl group is the most labile of those

$$o\text{-CH}_3\text{C}_6\text{H}_4\text{CH}_3\text{N}(\text{CH}_3)\text{CH}_2\text{C}_6\text{H}_4\text{Cl-}p \xrightarrow{\text{BrCN}}$$

p-ClC₆H₄CH₂N(CN)CH₃ + o-CH₃C₆H₄CH₂Br

studied; 23 no data are available permitting a direct comparison of it with the allvl group.

with the allyl group.

In a study of the relative ease of eleavage of amines containing substituted benzyl groups, von Braun and Engel ¹⁶ observed a close relationship between the ease of eleavage and the rate with which similarly substituted benzyl chlorides react with ethoxide ion. In the accompanying table are given some second-order rate constants for the companying table are given some second-order rate constants for the reaction of several benzyl chlorides with ethoxide ion as determined by the method of Franzen.⁴¹ The increase in ease of removal of these benzyl groups from an amine by eyanogen bromide parallels the increase in these rate eonstants.

⁴¹ Franzen, J. prakt. Chem., [2] 97, 61 (1918).

RELATIVE REACTIVITIES OF SOME BENZYL CHLORIDES WITH ETHOXIDE ION

Chloride	k_2
Benzyl	7.9 ± 0.3
p-Methylbenzyl	11.9 ± 0.3
p-Ethylbenzyl	14.9 ± 0.8
<i>p</i> -Phenylbenzyl	73.8 ± 0.2

Though the allyl group is more labile than the benzyl group, introduction of some labilizing groups into the para position of the benzyl group causes a greater increase in lability than introduction of the same groups into the γ position of the allyl group. This is shown by the accompanying reactions, ¹⁶ for which only the major products are given.

$$p\text{-RC}_6\text{H}_4\text{CH}_2\text{N}(\text{CH}_3)\text{CH}_2\text{CH} = \text{CHR} \xrightarrow{\text{BrCN}}$$

$$R = C_6\text{H}_5 \text{ or CH}_3$$

$$RCH = CHCH_2N(CN)CH_3 + p-RC_6H_4CH_2Br$$

The effect of structure on the ease of cleavage of various substituted allyl and benzyl groups is closely analogous to the effect on the reactivities of the corresponding allyl and benzyl halides in second-order displacement reactions. For example, those substituents that have been shown to increase the ease of cleavage of the benzyl group from an amine by cyanogen bromide also increase the reactivity of the benzyl halides in displacement reactions.

The Cyanomethyl Group. The ease of cleavage of the cyanomethyl group ⁴² has been estimated to be approximately equal to that of the ethyl group. Diethylaminoacetonitrile undergoes cleavage in both directions in nearly equal amounts. Similar behavior is exhibited by the

$$(C_2H_5)_2NCH_2CN \xrightarrow{BrCN} C_2H_5N(CN)CH_2CN + C_2H_5Br \\ \rightarrow (C_2H_5)_2NCN + BrCH_2CN$$

carbethoxymethyl group. Cleavage of dimethylaminoacetonitrile proceeds nearly completely in the direction yielding methyl bromide. The cyanomethyl group reduces the ease with which an amine will react with cyanogen bromide. When methylanilinoacetonitrile is treated with cyanogen bromide at 100° for five hours, bromination of the ring occurs in preference to cleavage of the amine.⁴³ No reaction takes place at room temperature.

$$C_6H_6N(CH_3)CH_2CN \xrightarrow{BrCN} p-BrC_6H_4N(CH_3)CH_2CN$$

⁴² von Braun, *Ber.*, 40, 3933 (1907). ⁴³ von Braun, *Ber.*, 41, 2100 (1908).

Methylenediamines. The methylenic linkage in tetrasubstituted methylenediamines is cleaved by cyanogen bromide with extreme ease. 44

$$[(C_6H_5CH_2)_2N]_2CH_2 \xrightarrow{2BrCN} 2(C_6H_5CH_2)_2NCN + CH_2Br_2$$

Even when the labile benzyl group is present, cleavage proceeds exclusively in the direction shown.³

A Steric Anomaly. A peculiar steric effect involving the reaction of some *ortho*-substituted aromatic amines has been observed. Some diphenylmethane derivatives containing two dimethylamino groups both of which are hindered by a group in the *ortho* position, e.g., I and II,

undergo no reaction with cyanogen bromide. Attributing this lack of reactivity to a steric or *ortho* effect, one would predict that compounds of a similar type containing one hindered and one unhindered dimethylamino group, e.g., III and IV, would react only at the unhindered group. However, under the same conditions these compounds react at both dimethylamino groups.⁴⁶ A similar situation has been observed when

$$(CH_3)_2N \longrightarrow CH_2 \longrightarrow N(CH_3)_2 \xrightarrow{2BrCN} N(CH_3)_2 \longrightarrow CH_3 \cap CH_3 \cap$$

these compounds were treated with iodoacetonitrile. Analogous compounds in the biphenyl series give the same results. No satisfactory explanation of this anomaly has been offered.

⁴⁴ von Braun and Röver, Ber., 36, 1196 (1903).

s von Braun and Kruber, Ber., 46, 3470 (1913).

⁴ von Braun and Mintz, Ber., 50, 1651 (1917).

CYCLIC AMINES

An aspect of the reaction of nitrogen ring compounds with cyanogen bromide that has received considerable study is the determination of the relative ease of fission of various ring systems. In the method most frequently employed, the ratios of ring cleavage to dealkylation of different rings containing the same alkyl group as a substituent on the nitrogen atom are compared. From a knowledge of the relative ease of displacement of several of the alkyl groups discussed previously, it is frequently possible to select a substituent that permits either complete dealkylation or complete cleavage of the ring.

Ethylenimines. Ethylenimines are known to undergo ring cleavage very readily in the presence of electrophilic reagents, i.e., compounds that convert the amino nitrogen to the quaternary state. Therefore, it is not surprising that this ring system is readily cleaved by cyanogen bromide. Only four examples of the reaction of 1-substituted ethylenimines with cyanogen bromide have been reported. By the gradual addition of 1-ethyl- or 1-n-butyl-ethylenimine to an ether solution of cyanogen bromide, there are obtained 88% and 94% yields, respectively, of the β-bromoethylcyanamides. The ring system in ethylenimines is so labile that it is doubtful if any substituent could be displaced from the

$$\begin{array}{ccc}
CH_2 & \xrightarrow{BrCN} & BrCH_2CH_2N(CN)R \\
N & & & \\
R & = C_2H_5 \text{ or } n\text{-}C_4H_9
\end{array}$$

nitrogen without cleaving the ring.

Cleavage of symmetrical rings of the type shown above can yield only one bromoalkyl cyanamide. An unsymmetrical cyclic structure offers the possibility of cleavage in two directions. Only a few examples of the unsymmetrical type have been reported. Three products were obtained from the reaction of 1-n-butyl-2-ethylethylenimine with cyanogen bromide in ether solution. This cleavage at the secondary alkyl linkage

$$\begin{array}{c} H_5C_2CH & \xrightarrow{CH_2} CH_2 & \xrightarrow{BrCN} & C_2H_5CHBrCH_2N(CN)C_4H_9-n \\ N & & & \\ C_4H_9-n & & & \end{array}$$

+
$$CH_3CH$$
= $CHCH_2N(CN)C_4H_9-n + C_2H_5CH(NHC_4H_9-n)CH_2Br \cdot HBr$
11%

rather than at the primary alkyl linkage is inconsistent with the greater ease of cleavage of the *n*-propyl group compared to the isopropyl group

(see p. 206) and the direction of cleavage of 1-n-butyl-2-methylpyrrolidine (see p. 214). The greater strain in the ethylenimine ring may account for this difference.

The reaction of 1-n-butyl-2,2-dimethylethylenimine 5 with cyanogen bromide yields a considerable quantity of an unidentified polymeric material. The only discrete products isolated are those shown in the

$$(CH_3)_2C$$
 CH_2
 \xrightarrow{BrCN}
 C_4H_9 - n

$$CH_2 = C(CH_3)CH_2N(CN)C_4H_9 - n + (CH_3)_2CBrCH_2NHC_4H_9 - n \cdot HBr_{29\%}$$

accompanying formulation. These results show that ring cleavage occurs preferentially at the tertiary alkyl linkage by an elimination reaction. The hydrogen bromide produced accounts for the observed formation of polymeric material.

Azetidines. The only azetidine whose reaction with cyanogen

bromide has been reported is 1-n-butylazetidine.⁵

$$\begin{array}{c} \mathrm{CH_2} \\ \mathrm{CH_2} \\ \mathrm{CH_2} \\ \end{array} \xrightarrow{\mathrm{BrCN}} \ \mathrm{Br(CH_2)_3N(CN)C_4H_9} - n \\ \mathrm{N} \\ \mathrm{C_4H_9} - n \end{array}$$

Pyrrolidines and Other Five-Membered Rings. Simple pyrrolidines are considerably more resistant to ring cleavage than are ethylenimines. Varying degrees of stability are observed in related compounds such as dihydroindoles, dihydroisoindoles, and indolizidines.

When treated with cyanogen bromide in benzene solution, 1-nbutylpyrrolidine gives a quantitative yield of n-butyl- δ -bromobutyleyanamide. 5.47 Even when the more labile ethyl group is employed as the

substituent, the ring is cleaved to the extent of 94%.48 However, when a benzyl group is employed as the substituent, the pyrrolidine ring is not

⁴ Ochiai, Tsuda, and Yokoyama, Ber., 68, 2291 (1935).

⁴⁸ von Braun, Ber., 44, 1252 (1911).

opened.49 A few unsymmetrical pyrrolidines undergo ring cleavage in 214

$$\begin{array}{c}
\text{A lew unsymmet} \\
\text{M} \\
\text{CH}_2\text{C}_6\text{H}_4\text{CH} = \text{CH}_2\text{-0} \\
\text{CN}
\end{array}$$

both possible directions. The ring opening of 1-n-butyl-2-methylpyrrolidine proceeds predominantly to yield the primary alkyl bromide.5

$$\begin{array}{c} & \xrightarrow{\text{BrCN}} \\ & \stackrel{N}{\underset{C_4H_9-n}{}} \end{array}$$

n
 Br(CH₂)₂CH(CH₃)N(CN)C₄H₉- n + CH₃CHBr(CH₂)₃N(CN)C₄H₉- n 26%

When the isopropyl group is present instead of the n-butyl group, cleavage still gives predominantly the primary bromide, but the 1-phenyl analog 50 cleaves to yield the secondary bromide as the major product.

Reaction of 1-n-butyl-2,2-dimethylpyrrolidine with cyanogen bromide proceeds exclusively by cleavage at the tertiary alkyl linkage.⁵ This

mode of cleavage, which is analogous to that of the similarly substituted ethylenimine (see p. 213), indicates that cyanogen bromide removes a tertiary alkyl group from an amine by an elimination reaction more readily than it removes a simple primary alkyl group by a displacement reaction. Compared with the pyrrolidine ring, the dihydroindole ring is slightly more susceptible to cleavage. 51

$$\begin{array}{c|c} & \xrightarrow{BrCN} & \xrightarrow{CH_2CH_2Br} + \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\$$

⁴⁹ von Braun, Ber., 49, 2629 (1916).

⁵⁰ Elderfield and Green, J. Org. Chem., 17, 431 (1952).

⁵¹ von Braun, Ber., 51, 96 (1918).

The ring system in dihydroisoindoles contains carbon-nitrogen bonds of the benzyl type; dihydroisoindoles are, accordingly, more susceptible to ring fission than dihydroindoles. The ring is sufficiently stable, however, to permit the removal of a benzyl group without cleavage of the ring, as shown by the accompanying equation.⁵²

When the substituent on the nitrogen of a dihydroisoindole is a methyl group, ring opening occurs more readily than demethylation.⁵³

Piperidines and Other Six-Membered Rings. A direct comparison of the relative stability of the piperidine and pyrrolidine rings is afforded by the reaction of indolizidine 54 with cyanogen bromide. The direction of ring cleavage was determined by degradation of the reaction product to racemic conline. Though 1-ethylpyrrolidine undergoes nearly

$$\begin{array}{c|c}
\hline
& BrCN \\
\hline
& N \\
\hline
& CH_2CH_2CH_2Br
\end{array}$$

complete ring eleavage, 1-ethylpiperidine undergoes de-ethylation to the extent of 66%. The ease of cleavage of the piperidine ring is roughly equal to the ease of removal of the n-propyl group as shown by the reaction of 1-n-propylpiperidine is with cyanogen bromide. Benzyl

groups can be removed with no detectable cleavage of the piperidine ring, 4.11

An excellent example of an elimination reaction is furnished by the behavior of ethyl β -(1-piperidyl)propionate.³ This is the only reported example of the reaction of a β -amino acid ester with cyanogen bromide.

$$\xrightarrow{\text{BrCN}} \xrightarrow{\text{BrCN}}$$

$$\xrightarrow{\text{CH}_2\text{CH}_2\text{CO}_2\text{C}_2\text{H}_5} + \xrightarrow{\text{CH}_2\text{CH}_2\text{CO}_2\text{C}_2\text{H}_5} + \text{CH}_2 = \text{CHCO}_2\text{C}_2\text{H}_5$$

$$\xrightarrow{\text{N}} \xrightarrow{\text{CH}_2\text{CH}_2\text{CO}_2\text{C}_2\text{H}_5} \cdot \text{HBr} \quad \text{CN}$$

To insure that cleavage of the piperidine ring will be predominant, the substituent should possess a resistance to cleavage equal to or greater than that of the *n*-butyl group. Surprisingly, 1-isopropyl-4-pipecoline is reported ⁵⁶ to undergo dealkylation with no detectable ring cleavage. The reaction of 1-phenylpiperidine ^{7,57} can result only in ring opening since the phenyl group cannot be displaced.

Tropane, which contains both the piperidine and pyrrolidine ring systems, is completely demethylated by cyanogen bromide. Under the

$$NCH_3 \xrightarrow{BrCN} NCN + \left[N(CH_3)_2 \right]$$
Br

conditions employed for this reaction, nearly half the tropane was converted to the quaternary salt by reaction with the methyl bromide formed.⁴⁸

Tetrahydroquinoline is slightly more resistant to ring cleavage than piperidine. For 1-n-propylpiperidine ⁴⁸ the ratio of ring opening to depropylation is 3:2; for 1-n-propyltetrahydroquinoline ⁵⁸ this ratio is 3:4. The contrasting modes of reaction of 1-methyl-3-phenyltetrahydroquinoline and 1-methyl-2-phenyltetrahydroquinoline show how the stability of the ring can be modified. From the former the only product isolated was 1-cyano-3-phenyltetrahydroquinoline, whereas from the latter there resulted a 50% yield of the ring-opened product.⁵⁹

⁵⁸ Elderfield, Pitt, and Wempen, J. Am. Chem. Soc., 72, 1344 (1950).

⁵⁷ von Braun, Ber., 41, 2165 (1908). ⁵⁸ von Braun, Ber., 42, 2219 (1909).

¹⁹ von Braun, Seemann, and Schultheis, *Ber.*, 55, 3803 (1922).

goes ring opening with no appreciable demethylation, the o-ethylbenzyl group is removed in preference to cleavage of the ring.⁶²

$$\begin{array}{c}
O \\
\stackrel{\text{BrCN}}{\longrightarrow} o\text{-C}_2\text{H}_5\text{C}_6\text{H}_4\text{CH}_2\text{Br} + O \\
N \\
\text{CH}_2\text{C}_6\text{H}_4\text{C}_2\text{H}_6\text{-}o
\end{array}$$

In benzomorpholine the ring is considerably more stable than in morpholine. Reaction of 4-methylbenzomorpholine with cyanogen bromide results in recovery of half of the starting material; no product

$$\begin{array}{c|c}
O & \xrightarrow{BrCN} & O \\
& & \\
CH_3 & & \\
\end{array}$$

$$\begin{array}{c}
O \\
+ \\
CN & \\
\end{array}$$

$$\begin{array}{c}
O \\
N \\
(CH_3)_2
\end{array}$$

$$\begin{array}{c}
Br \\
\end{array}$$

resulting from ring opening is obtained.63

The piperazine ring is the most readily cleaved of the six-membered rings that have been studied. When cyanogen bromide is added to 1,4-dimethylpiperazine, the major products isolated are the hydro-

$$2H_3CN$$
 $NCH_3 \xrightarrow{B_1CN}$
 H_3CN
 $NCH_3 \cdot 2HBr + 2CH_2 = CHN(CN)CH_3$

bromide of the starting material and methylvinyleyanamide.64

Pyridines and Quinolines. Reaction of γ -dipyridyl in absolute ethanol with two moles of cyanogen bromide gives an adduct whose composition corresponds to the addition of one mole of cyanogen bromide. This is one of the few adducts of this type to have been isolated

$$X \xrightarrow{2BrCN} X \xrightarrow{Br} XCX$$

and characterized. Reaction of pyridine with cyanogen bromide, followed by treatment with a primary or secondary amine, gives products

believed to result from the intermediate formation of 1-cyano-2-bromo-Quinoline reacts with cyanogen bromide in 1,2-dihydropyridine.66

$$\begin{array}{c|c} \text{H} & \text{H}_{20} \\ \hline \\ \text{CN} & \text{Br} \\ \hline \\ \text{CN} & \text{OH} \\ \end{array}$$

moist ether to give 1-cyano-2-hydroxy-1,2-dihydroquinoline and its ether. 67, 68, 69

Simultaneous reaction of quinoline with cyanogen bromide and anhydrous hydrogen cyanide in benzene at 0° yields 1,2-dicyano-1,2-di-

2
$$+ BrCN + HCN \rightarrow$$
 $CN + HCN + HCN \rightarrow$
 $CN + HCN + HCN \rightarrow$
 $CN + HCN + HCN \rightarrow$
 $CN +$

hydroquinoline. 68,70 If the quinoline ring contains substituents in the 2 or 8 position, this reaction takes place less readily and it is necessary to operate in sealed tubes at 150°. The structures of these products

were established by conversion to the quinolinecarboxylic acids.

⁶⁶ Migrdichian, The Chemistry of Organic Cyanogen Compounds, p. 110, Reinhold Publishing Corp., New York, 1947.

⁵ Shimidzu, J. Pharm. Soc. Japan, 529, 243 (1926) [C. A., 20, 2680 (1926)].

⁶³ Mumm and Ludwig, Ann., 514, 34 (1934).

⁶⁹ von Braun, Wallach-Festschrift, 313 [C. A., 5, 888 (1911)].

⁷⁰ Mumm and Herrendorfer, Ber., 47, 75S (1914).

Alkaloids

The von Braun cyanogen bromide reaction has frequently been employed in the degradation of alkaloids by attack at the basic nitrogen atoms. Its importance in this field is comparable to that of the classical Hofmann and Emde methods of degradation. Another reaction bearing von Braun's name, which also has found considerable application as a method of degradation, consists in dealkylation of secondary amines by preparing the benzoyl derivative and treating this amide with phosphorus pentachloride or bromide.

A few examples of the reaction of cyanogen bromide with alkaloids are presented merely to indicate the applicability of the reaction in this field. No detailed coverage or critical evaluation in relation to other methods of degradation ⁷¹ is intended.

The value of any reaction to be used as a method of degrading compounds of unknown structure is greatly enhanced by a thorough understanding of the course of the reaction when applied to many simple compounds of known structure. The examples discussed above have aided in the development of this reaction as a method of degradation.

Though repeated application of Hofmann's method of exhaustive methylation often effects complete removal of a nitrogen atom, originally part of a heterocyclic ring, this cannot be accomplished by the use of cyanogen bromide. On the other hand, cyanogen bromide will sometimes effect ring opening where the Hofmann method fails, namely, in the dihydroindole and tetrahydroquinoline ring systems.⁴⁹ Hydrocotarnine (VI) provides an example of the degradation of a compound in different ways by the Hofmann and von Braun methods.^{61,72} This cxample also illustrates some of the deductions that can be made from the reaction of a compound with cyanogen bromide. Analysis of the

$$\begin{array}{c|c} O & & & & & \\ \hline O & & & & \\ \hline CH_2 & & & & \\ \hline O & & & & \\ \hline CH_2 & & & \\ \hline O & & \\ \hline CH_2 & & \\ \hline O & & \\ \hline CH_2 & & \\ \hline O & & \\ \hline CH_2 & & \\ \hline CH_2 & & \\ \hline O & & \\ O & & \\ \hline O & & \\ O & & \\ \hline O & & \\ O & & \\ \hline O & & \\ O & & \\ \hline O & & \\ O & & \\ \hline O & & \\ O & & \\ \hline O &$$

ⁿ Houben, Die Methoden der organischen Chemie, 2nd ed., Vol. IV, pp. 519-526, G. Thieme, Leipzig. 1924.

"Small, in Gilman, Organic Chemistry, Vol. II, 2nd ed., p. 1175, John Wiley & Sons, New York, 1943. reaction product VII, showing that the elements of cyanogen bromide have been added, implies that a tertiary amine nitrogen atom constitutes part of a ring that has undergone opening. Once the presence of an N-methyl group has been established, it can be concluded that the nitrogen ring system is one that is sufficiently labile to undergo ring cleavage in preference to demethylation. This indicates that a stable ring of the piperidine or tetrahydroquinoline type is probably not involved. The observed behavior, however, is compatible with ring systems such as dihydroindole, dihydroisoindole, or tetrahydroisoquinoline. A selection among these possibilities will be dictated by other consistent experimental data.

Conessine (VIII), whose structure is not known, reacts with one equivalent of cyanogen bromide in ether solution to give two principal products. One of these (IX), which proved to be a quaternary ammonium salt, is doubtless formed by the reaction of two moles of methyl

It, is doubtless formed by one real
$$C_{24}H_{40}N_2 + BrCN \rightarrow C_{26}H_{46}N_2Br_2 + C_{23}H_{37}N_2CN$$

VIII

(Y) has the $C_{24}H_{40}N_2 + BrCN \rightarrow C_{26}H_{46}N_2Br_2 + C_{23}H_{37}N_2CN$

bromide with the starting material. The other (X) has the composition of a cyanamide arising from a demethylation of conessine. Further treatment of the cyanamide X with cyanogen bromide yields a product

$$\begin{array}{c} \text{C}_{23}\text{H}_{37}\text{N}_2\text{CN} + \text{BrCN} \rightarrow \text{C}_{22}\text{H}_{34}\text{N}_2(\text{CN})_2 \\ \text{X} \end{array}$$

(XI) arising from a second demethylation. These results strongly indicate that each of the nitrogen atoms in conessine contains at least one methyl group. Furthermore, these amine functions much be joined to the molecule by bonds more stable with respect to charge by cyanogen bromide than the N-methyl bond.

An interesting application of the cyanogen bromide reaction to the morphine alkaloids is the comparison of the behavior of diagratylmorphine (XIV), which undergoes demethylation, with that of the behavior (XIV), which adds the elements of cyanogen bromide.⁷⁴

CH₃ CH₂

$$CH_2$$

$$CH_2$$

$$CH_2$$

$$CH_3$$

$$CH$$

Siddiqui and Siddiqui, J. Indian Chem. Soc., 11, 787 (1934).
 von Braun, Kruber, and Aust, Ber., 47, 2312 (1914).

The only pertinent structural difference in the nitrogen ring system of these two compounds is the presence of β,γ unsaturation between carbon atoms 8 and 14 in thebaine (XIV) in contrast to the more remote γ,δ unsaturation at the 7-8 position in diacetylmorphine (XII). The β,γ -double bond in position 8-14 involves an allylic linkage to the nitrogen atom which labilizes the nitrogen ring system to a considerable extent. This explanation is supported by the fact that tetrahydrothebaine undergoes demethylation rather than ring cleavage. Demethylation rather than ring cleavage of morphine and codeine is one reason for assigning the double bond in these compounds to position 7-8 rather than to position 8-14.

When optically active dibenzoylapomorphine (XVI) is treated with eyanogen bromide in ehloroform solution, there is obtained a 50% yield of a product resulting from ring opening and simultaneous loss of hydrogen bromide.⁷⁵ Though the analytical figures obtained for the

$$C_{\epsilon}H_{5}COO \ OCOC_{\epsilon}H_{\delta}$$
 $C_{\epsilon}H_{5}COO \ OCOC_{\epsilon}H_{\delta}$
 $C_{\epsilon}H_{5}COO \ OCOC_{\epsilon}H_{\delta}$
 $C_{\epsilon}H_{5}COO \ OCOC_{\epsilon}H_{\delta}$

product are equally satisfactory for a compound arising from demethylation without ring opening, structure (XVII) is assigned on the basis of the observed loss in optical activity. Furthermore, the course of the reaction as indicated is consistent with the known lability of a benzyl linkage.

In connection with the problem of the determination of the structure of lupinine. Winterfeld and Holschneider have treated lupinane (XVIII) with cyanogen bromide in boiling benzene. Occurrence of the

[&]quot; son Braun and Aud, Ber., 50, 43 (1917).

Winterfeld and Holschneider, Ben., 64, 137 (1931).

ring cleavage predominantly in the direction indicated, rather than with fission of the other ring, was demonstrated by degradation of the

$$\begin{array}{c|c} CH_3 & \xrightarrow{BrCN} & CH_3 \\ \hline N & (CH_2)_4 Br \end{array} \rightarrow \begin{array}{c|c} CO_2 H \\ \hline N & CO_2 H \end{array}$$

product (XIX) to quinolinic acid (XX). Had ring cleavage in the reverse direction predominated, the ultimate product would have been α -picolinic acid.

Sparteine (XXI) reacts with cyanogen bromide 77 to yield three ring-

opened products, one resulting from the addition of two moles of cyanogen bromide and two incorporating one mole of cyanogen bromide, whose structures have not been determined.

When treated with cyanogen bromide in chloroform solution, cocaine (XXII) undergoes ring opening to only a very slight extent; demethylation is the predominant reaction. Some cocaine methobromide results from reaction of the liberated methyl bromide with cocaine.

$$\begin{array}{c|c} \operatorname{CH_3O_2C} & \operatorname{CH_3O_2C} \\ \operatorname{C_6H_5CO_2} & \operatorname{NCH_3} \xrightarrow{\operatorname{BrCN}} \operatorname{C_6H_5CO_2} & \operatorname{NCN} + \operatorname{CH_3Br} \end{array}$$

Treatment of the reaction product (XXIII) with concentrated hydrochloric acid at 120° causes the elimination of benzoic acid and removal of the cyano group, thereby yielding desmethylanhydroecgonine. The ethyl ester of anhydroecgonine (XXIV) cannot be demethylated by cyanogen bromide in an appreciable yield because of extensive ring cleavage. The enhanced lability of the ring in XXIV can be attributed to the presence of β,γ unsaturation.

⁷⁷ Winterfeld and Holschneider, Arch. Pharm., 267, 433 (1929).

⁷⁸ von Braun and Müller, Ber., 51, 235 (1918).

SYNTHETIC APPLICATIONS

Occasional mention of the synthetic value of the von Braun cyanogen bromide reaction can be found in the literature.^{3, 5, 55, 57, 79, 80} The adoption of this reaction for large-scale synthesis is limited by the properties of cyanogen bromide; its toxicity and volatility discourage the handling of large quantities of cyanogen bromide. The instability of cyanogen bromide makes it inadvisable to attempt to store large quantities of it for an indefinite period. Consequently, use of the cyanogen bromide reaction in synthesis is at present restricted to the field of rare chemicals. The following survey of some applications, together with a few suggested uses, is intended to provide an evaluation of the potentialities of the reaction in syntheses.

The preparation of alkyl bromides by the cleavage of acyclic amines with cyanogen bromide finds only limited use, since these bromides are obtained more readily by other methods. However, the cyanogen bromide reaction does provide a convenient synthesis of bromoacetonitrile (p. 228) and of o-vinylbenzyl bromide (p. 228).

The alkylation of eyanamide frequently offers a convenient synthesis of dialkylcyanamides containing two identical substituents, but this method is of little value when two different substituents are desired. The direct introduction of an aryl group into cyanamide is also not readily accomplished. To obtain a cyanamide containing one aryl and one alkyl group, it is often possible to remove one alkyl group from a dialkylarylamine by treatment with eyanogen bromide. Cressman so has employed the cyanogen bromide reaction for the preparation of monoalkyl α -naphthylcyanamides from dialkyl α -naphthylamines. The hydrolysis of unsymmetrically substituted cyanamides offers a means of obtaining unsymmetrical secondary amines in a pure state. Since guanidines are readily prepared by the reaction of cyanamides with amine salts, the applicability of the eyanogen bromide reaction to the synthesis of unsymmetrically substituted guanidines is apparent.

$$\begin{array}{c} R' \\ \text{NCN} + R'' \text{NHR'''} \cdot \text{HX} \rightarrow \\ R \end{array} \xrightarrow{R'} \begin{array}{c} \text{NH} \quad R'' \\ \text{NGN} \end{array} \cdot \text{HX}$$

The bromoalkylcyanamides obtained by ring cleavage are more useful since they can be employed in the synthesis of compounds that

⁷ von Braun, Ber., 41, 2113 (1908).

^{**} Cressman, Org. Syntheses, 27, 56 (1947).
** Erlenmeyer, Ann., 146, 258 (1868).

frequently are difficult to obtain by other methods. The β -bromoethylalkylcyanamides resulting from the ring opening of 1-alkylethylenimines react with primary amines to yield various cyclic guanidine derivatives and with secondary amines to give, after hydrolysis, unsymmetrical derivatives of ethylenediamine.⁵ The products obtained by the

$$\begin{array}{c} CH_2 - CH_2 \\ BrCH_2CH_2N(CN)R + R'NH_2 \rightarrow RN & NR' \cdot HBr \\ C & \parallel \\ NH \\ \\ BrCH_2CH_2N(CN)R + R'NHR'' \rightarrow & NCH_2CH_2N(CN)R \\ R' & \downarrow H_2O \\ R'' & NCH_2CH_2NHR \\ R' & \end{array}$$

ring opening of 1-alkylpyrrolidines have served as intermediates for the preparation of derivatives of putrescine ⁵ and monoalkylamino derivatives

$$Br(CH_2)_4N(CN)R + R'NHR'' \rightarrow R'R''N(CH_2)_4N(CN)R$$

$$\downarrow H_2O$$

$$R'R''N(CH_2)_4NHR$$

of valeric acid.47 The product from the cleavage of N-phenylpiperidine

$$RN(CN)(CH_2)_4Br \xrightarrow{(2) \text{ Hydrolysis}} RNH(CH_2)_4COOH$$

with cyanogen bromide has been used for the synthesis of N,N'-diphenyl-cadaverine. 57

The above examples illustrate some applications of bromoalkyleyanamides to the synthesis of compounds through replacement of the bromine atom by nucleophilic reagents without altering the cyanamide portion of the molecule. Though the recorded examples of the use of these bromoalkyleyanamides are few, they suggest a wide variety of applications to be investigated.

EXPERIMENTAL CONDITIONS AND PROCEDURES

Solvents. Many procedures in the literature describe the reaction of amines with eyanogen bromide in the absence of a solvent. This practice frequently gives poor yields because of unfavorable side reactions. Particularly for amines that react vigorously with eyanogen bromide, the use of a diluent is necessary to keep the reaction under control. With the less reactive derivatives of aromatic amines a solvent is less essential and has frequently been omitted. The omission of a solvent appears to offer little or no advantage. If a reaction requires heating, the selection of a solvent having an appropriate boiling point affords a simple means of maintaining adequate temperature control. The physical properties of cyanogen bromide are such (m.p. 52°; b.p. 62°) that heating a reaction mixture containing no solvent occasionally results in a clogged condenser. The use of a solvent accompanied by stirring gives more intimate mixing and avoids excessive local heating.

Non-polar solvents such as ether, chloroform, benzene, and the hydroearbons are to be preferred because of their immiscibility with water and their tendency to precipitate such by-products as amine salts, which can then be removed by filtration. Dry dioxane is a suitable solvent for the reaction but is to be avoided, if possible, since its miscibility with water complicates working up the reaction mixture. Though glacial acetic acid has been used, ⁵² hydroxylated solvents are generally less desirable. Reasonably anhydrous conditions are recommended to avoid interference associated with formation of hydrobromic acid.

Order of Mixing Reactants. An important factor is the order of addition of the reactants. As a general rule, the gradual addition of a solution of amine to a solution of eyanogen bromide is preferred. The reasons for this preference become evident when the predominant side reactions are considered. When highly reactive bromides such as allyl, benzyl, and methyl bromide are formed in the reaction, the presence of excess amine is conducive to the formation of quaternary ammonium bromides. Usually eyanogen bromide reacts with an amine more rapidly than do alkyl bromides, and use of the recommended order of addition minimizes this side reaction. Since hydrogen bromide reacts more rapidly with amines than does cyanogen bromide, the order of addition in elimination reactions in which hydrogen bromide is formed is of relatively little importance. Here the yields of olefin and disubstituted cyanamide are limited to a maximum of 50%, regardless of the order of addition.

If an amine is not very reactive toward cyanogen bromide, it will probably not react rapidly with an alkyl bromide. For such amines the simplest procedure is to mix the amine and cyanogen bromide in an appropriate solvent and then heat for the required time. Unless warranted by some special circumstance, such as the desire to cleave an amine in the presence of a thio ether group or to bring about preferential reaction of one of two amine functions present in the same molecule, the gradual addition of cyanogen bromide to an amine should be avoided.

With sensitive amines such as the ethylenimines it is almost imperative that the recommended order of addition be followed, since these amines tend to undergo extensive polymerization initiated by traces of a reactive alkyl halide or an acid.⁸³

Isolation of Products. Procedures for the reaction of an amine with cyanogen bromide are generally simple and not subject to wide variation. A greater variety of procedures is involved in working up the reaction mixture and in the isolation of a particular reaction product. The amine and evanogen bromide are allowed to react either without a solvent or, more frequently, in an inert, water-immiscible solvent such as ether. benzene, or chloroform. After completion of the reaction the addition of more solvent precipitates the major part of any quaternary ammonium salt or amine hydrobromide formed as by-products. Extraction of the solution with dilute aqueous acid removes any unreacted amine and the last traces of salts. The alkyl bromide and the cyanamide remaining in the organic layer can frequently be separated by fractional distillation. If distillation or crystallization does not effect a separation, the choice of another method depends upon whether the alkyl bromide or the cyanamide is the preferred product. By refluxing the mixture with hydrobromic acid it is often possible to hydrolyze the cyanamide to the amine hydrobromide and then isolate the desired alkyl bromide by steam distillation or extraction. If a particular derivative of the alkyl bromide is sought, it is often possible to carry out the reaction involving the alkyl bromide in the presence of the contaminating cyanamide and then to separate the derivative from the cyanamide. More frequently the cyanamide is the desired product. In such cases the contaminating alkyl bromide can be removed readily by reaction with a secondary or tertiary amine, followed by a separation of the amine salts from the neutral cyanamide.

These methods are generally applicable to cyclic as well as to acyclic amines. A paper by von Braun ³ is of particular interest in regard to the

⁸³ Fruton, in Elderfield, Heterocyclic Compounds, Vol. 1, p. 70, John Wiley & Sons, New York, 1950; Lassell and Sundet, J. Am. Chem. Soc., 63, 2374 (1941).

use of different methods for separating the products resulting from the reaction of several piperidine derivatives with cyanogen bromide.

Preparation and Properties of Cyanogen Bromide. A convenient preparation of cyanogen bromide in 200-300-g. quantities and in 73-85% yield from bromine and sodium cyanide is described in Organic Syntheses. In contrast to a note in this procedure on the instability of cyanogen bromide, the author has found that no decomposition occurred after storing in a glass-stoppered flask at room temperature for as long as a month. The toxicity and volatility of cyanogen bromide require that all operations with this material be performed in an efficient hood.

The cleavage of dimethyl- α -naphthylamine with cyanogen bromide to furnish methyl- α -naphthylcyanamide in 63-67% yield is described in Organic Syntheses.⁸⁰

Bromoacetonitrile.⁷⁹ When 200 g.* (1.61 moles) of N-eyanomethyl-piperidine is mixed with 171 g. (1.61 moles) of cyanogen bromide, an exothermic reaction occurs, accompanied by the formation of a solid. After the reaction has subsided, the mixture is allowed to stand overnight. Though the reaction is essentially complete at this stage, the mixture is heated for a short time on the steam bath. This heating removes the greater part of any unreacted cyanogen bromide. Ether is added to the cooled reaction mixture, and the solid (quaternary salt formed by reaction of 1-cyanomethylpiperidine with bromoacetonitrile) is removed by filtration. The ether solution is extracted with water to remove the last traces of the quaternary salt, the solvent is removed, and the residual yellow oil is vacuum distilled. There is obtained 135–140 g. (about 70%) of bromoacetonitrile collected over the range 50–90°/15 mm., the greater part distilling at 50°. The residual N-cyanopiperidine distills at 115°/15 mm.

The crude bromoacetonitrile is pure enough for most purposes. A second distillation gives the pure product, a strongly lachrymatory liquid, b.p.46°/13 mm. or 150-151°/752 mm.

o-Vinylbenzyl Bromide. Treatment of an ice-cold ether solution of o-vinylbenzyldimethylamine with eyanogen bromide causes the precipitation of di-(o-vinylbenzyl)dimethylammonium bromide, m.p. 178-179°. After filtration, the ether solution containing the o-vinylbenzyl bromide and dimethyleyanamide is extracted with dilute aqueous acid to remove unchanged amine and the water-soluble dimethyleyanamide. After drying the ether solution over calcium chloride and removing the

¹⁴ Hartman and Dreger, Org. Syntheses, Coll. Vol. II, p. 150, John Wiley & Sons, New York, 1941.

When small amounts of materials are used, the heat evolved is insufficient to cause
an appreciable reaction. The mixture is heated on the steam bath for two to three hours
in a scaled tube.

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ether, a colorless oil remains. Distillation gives colorless, analytically pure o-vinylbenzyl bromide, b.p. 119-120°/17 mm., in 50% yield.

n-Butyl- β -bromoethylcyanamide.⁵ A solution of 65 g. (0.65 mole) of 1-n-butylethylenimine in 300 ml. of absolute ether is added during four hours with stirring to a solution of 75 g. (0.71 mole) of cyanogen bromide in 200 ml. of ether. The heat of reaction is sufficient to maintain gentle refluxing of the ether. The mixture is allowed to stand overnight, and the clear, pale yellow ether solution is extracted with 100 ml. of 5% hydrochloric acid and two 100-ml. portions of water and then dried over calcium chloride. Removal of the ether and distillation of the residue (131 g.) gives 126 g. (94%) of n-butyl- β -bromoethylcyanamide as a colorless liquid, b.p. 106–108°/0.6 mm.

n-Butyl-4-bromopentylcyanamide and n-Butyl-(1-methyl-4-diethyl-aminobutyl)cyanamide.⁵ Addition over a four-hour period of a solution of 70.5 g. (0.50 mole) of 1-n-butyl-2-methylpyrrolidine in 200 ml. of benzene to a stirred solution of 58.2 g. (0.55 mole) of cyanogen bromide in 200 ml. of benzene gives a clear, pale yellow solution which is allowed to stand overnight. The benzene solution is extracted with 100 ml. of 5% hydrochloric acid and with two 100-ml. portions of water and dried over calcium chloride. Removal of the benzene under reduced pressure leaves 120 g. of a clear red-brown liquid. The theoretical yield of ring-opened product is 123 g.

This crude product (a mixture of isomers) is refluxed for three and one-half hours with 292 g. (4.0 moles) of diethylamine. After removal of excess diethylamine by distillation, the residue is treated with a solution of 50 ml. of concentrated hydrochloric acid in 200 ml. of water. The acid-insoluble oil is taken up in 350 ml. of ether and dried over calcium chloride. Removal of the ether leaves 32 g. of n-butyl-4-bromopentylcyanamide as a yellow liquid.

The hydrochloric acid extract is made strongly basic with potassium hydroxide. The oil that separates is taken up in 400 ml. of ether and dried over potassium carbonate. Removal of the ether and traces of diethylamine leaves 81 g. of a clear red-brown liquid. Distillation of 41 g. of this crude basic product gives 36 g. of n-butyl-(1-methyl-4-diethylamino-butyl)cyanamide as a pale yellow oil, b.p. 130-133°/0.7 mm.

Cyanonorcocaine. Cyanogen bromide (30 g.) is added to a solution of 100 g. of cocaine in 200 ml. of chloroform and the mixture refluxed on the steam bath for two hours. After removal of the chloroform the solid residue is treated with water. From the water solution there is obtained 8 to 9 g. of crude cocaine methobromide. One recrystallization of the water-insoluble solid from ethanol containing a little water gives 62–65 g. (60–63%) of pure cyanonorcocaine, m.p. 123–124°.

RELATIVE EASE OF CLEAVAGE OF AMINES BY CYANOGEN BROMIDE

No accurate tabulation of the relative lability of the various alkyl groups in respect to cleavage from amine nitrogen by cyanogen bromide can be constructed on the basis of the experimental work recorded in the literature.

Table I provides a general picture of the relative lability of the majority of the groups that have been studied. References concerning the groups listed in Table I are not included because an intricate system of cross references would be required. An amine containing a particular alkyl group listed in Table I can be located in Table III where it is accompanied by a literature reference. To emphasize the relation between some general classes of alkyl groups, the table has been arranged Column A contains groups of the allyl type, the in three columns. greater number of which have been compared directly with the unsubstituted allyl group. Column B is similarly arranged on the basis of the benzyl group; Column C with reference to the methyl group. The table is arranged in order of decreasing ease of removal of the group by cyanogen bromide. If two groups are widely separated vertically in the table, one can be reasonably sure that the group higher in the table will be cleaved much more readily than the lower member.

An evaluation of the relative lability of the rings in various cyclic amines can be made with more certainty than the relative lability of the alkyl groups mentioned above. By determining the ratio of ring opening to dealkylation of a particular cyclic amine as the substituents on the nitrogen are varied, a satisfactory estimation of the lability of the ring can be obtained. Though no quantitative conclusions are justified, the ring systems in Table II can be arranged on the basis of their relative lability with reasonable qualitative accuracy. The order of lability given is applicable only to the simple ring systems containing no activating or deactivating substituents in the ring. For example, a phenyl group in the 2 position of tetrahydroquinoline will cause this ring system to be more labile than the pyrrolidine ring. A few of the more pertinent references dealing with the ring systems listed are included.

TABLE I

RELATIVE EASE OF REMOVAL OF ALKYL GROUPS

(Descending in Order of Decreasing Lability)

 Λ

В

 \mathbf{C}

Methylene (diamines)

p-Methoxybenzyl

[p-Phenyl, p-cyclohexyl, and
p-xenylbenzyl] *
p-Ethylbenzyl

p-Methylbenzyl

γ-Phenylallyl γ-Ethylallyl γ-Methylallyl

Allyl

 α -Thienyl α -Furomethyl

2-Cyclopentenyl

m-Methyl- and o-phenyl-benzyl

 $[Benzyl \ {\it and} \ o ext{-}, m ext{-}, p ext{-} {\it fluorobenzyl}]$

 α -Naphthylmethyl β -Naphthylmethyl

[\gamma-n-Amylpropargyl, propargyl, and eyclopropylmethyl]

γ-Chloroallyl γ-Bromoallyl

β-Chloroallyl β-Bromoallyl

p-Chlorobenzyl

p-Bromo- and m-chloro-benzyl p-Iodo- and p-acetamido-benzyl m-Bromo- and m-acetamido-

benzyl m-Iodobenzyl

o-Chloro- and o-acetamido-benzyl

o-Bromobenzyl o-Iodobenzyl

p-Cyanobenzyl
o- and m-Cyanobenzyl
o-, m- and p-Nitrobenzyl

Methyl

[Ethyl, cyanomethyl, and carbalkoxy-methyl]
[Cyclobutylmethyl and n-propyl]
Phenethyl γ-Phenylpropyl
Isopropyl and n-butyl n-Amyl and isoamyl
[Isobutyl, n-hexyl and higher homologs]

^{*} Groups within brackets are of equivalent lability.

TABLE II

RELATIVE EASE OF RING CLEAVAGE OF CYCLIC AMINES

Amines Descending in Order of Decreasing Ease of Cleavage References

Note: References 85-112 are listed on p. 262.

TABULAR SURVEY

Tables III, IV, and V contain most of the known examples of the reaction of tertiary amines with cyanogen bromide involving the reaction discussed in this chapter. Particularly with respect to the alkaloids, the coverage is incomplete since a direct reference to the use of cyanogen bromide is often lacking. The literature has been covered through the year 1950.

Only the major products are listed in the tables. Where yields are available they appear in parentheses next to the product concerned. In several instances in which alkaloids were treated with cyanogen

bromide, either no structures or incorrect structures of the products were reported. Where correct structures are now available, these have been given rather than those reported in the reference cited.

The acyclic amines are covered in Table III, which is divided into the following sections: (A) Miscellaneous Aliphatic Amines; (B) Derivatives of Allylamine; (C) Derivatives of Benzylamine; (D) Derivatives of Other Arylmethylamines; (E) Derivatives of Aromatic Amines. Amines containing both the allyl and the benzyl groups are listed under Derivatives of Allylamine. Aromatic amines that contain the allyl and benzyl groups are listed under Derivatives of Aromatic Amines.

Table IV contains all cyclic amines except the alkaloids. It is divided into the following sections: (A) Three- and Four-Membered Rings (ethylenimines and azetidines); (B) Five-Membered Rings (pyrrolidines, dihydroindoles, and dihydroisoindoles); (C) Six- and Seven-Membered Rings (including piperidines, tetrahydroquinolines, morpholines, and piperazines). Bicyclic amines containing both five- and six-membered rings are included in this section. (D) Pyridine-Type Amines. Most of the examples in section D involve reactions of pyridines, quinolines, and related compounds with cyanogen bromide in which cyanogen bromide is considered to add across the 1,2 double bond to yield a 2-bromo-1-cyano-1,2-dihydro derivative. Occasionally the presence of nuclear substituents causes the cyanogen bromide to add 1,4 (see p. 219).

In Table V are listed most of the alkaloids whose reactions with cyanogen bromide are reported in the literature. Where the course of the reaction and the structure of the products are not known, only the empirical formulas are given.

In Table V and within the various sections of Tables III and IV the amines are listed in order of increasing number of carbon atoms.

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TABLE	

1		ORGANIC	REACTIONS		
Refer-	ence	444 42 1 1 1 442 1 1 1 1	442 37 37 37 37	42 44 37	39
ACYCLIC AMINES	Products Ational answers Attachetic Amines	CH ₃ N(CN)CH ₂ CN + $\{(CH_3)_3NCH_2CN]Br$ (ca. 50%) CH ₃ N(CN)CH ₂ CN + CH_2Br_2 C ₂ H ₆ N(CN)CH ₂ CN (40%) + BrCH ₂ CN (50%) (n-C ₃ H ₇) ₂ NCN + CH_3Br CH ₂ =CH(CH ₂) ₃ N(CN)CH ₃ + $\{CH_2$ =CH(CH ₂) ₃ N(CH ₃) ₃]Br (n-C ₃ H ₇) ₂ NCN + C ₂ H ₆ Br (n-C ₃ H ₇) ₂ NCN + n-C ₃ H ₇ N(CN)CH ₂ CO ₂ C ₂ H ₅ (C ₃ H ₅) ₂ NCN + r C ₂ H ₆ N(CN)CH ₂ CO ₂ C ₂ H ₅ (n-C ₃ H ₇) ₂ NCN + n-C ₃ H ₇ N(CN)CH ₂ CO ₂ C ₂ H ₅	[(CH ₃) ₂ CHCH ₂] ₂ NCN + BrCH ₂ CN n-C ₃ H ₇ N(CN)CH ₂ CO ₂ C ₂ H ₅ + (n-C ₃ H ₇) ₂ NCN C ₆ H ₆ (CH ₂) ₂ N(CN)CH ₃ + [C ₆ H ₆ (CH ₂) ₂ N(CH ₃) ₃]Br (40%) C ₆ H ₆ (CH ₂) ₃ N(CN)CH ₃ + [C ₆ H ₆ (CH ₂) ₃ N(CH ₃) ₃]Br (33%) No products isolated C ₆ H ₆ (CH ₂) ₂ N(CN)C ₂ H ₆ (70%) + C ₂ H ₆ Br C ₆ H ₆ (CH ₂) ₂ N(CN)C ₂ H ₆ (75%) + (C ₂ H ₆) ₂ NCN (25%) + C ₇ H ₇ (CH ₃) ₃ N ₅ .	No products isolated $(n\text{-}C_3H_7)_2 \text{NO} + \text{CH}_2 \text{Br}_2 + n\text{-}C_3H_7)_2 \text{NO} + \text{CH}_2 \text{Br}_2 + n\text{-}C_3H_7 \text{Br}_3 + n\text{-}C_3H_7 \text{Br}_4 + n\text{-}C_3H_7)_2 \text{NO} (35\%) + C_6 \text{H}_6 (\text{CH}_2)_3 \text{Br} (35\%) + C_6 \text{H}_6 (\text{CH}_2)_3 \text{Br} (35\%)$	D. Delivatives of Augments: $CH_2 = CCICH_2N(CN)CH_3 + CH_2 = CCICH_2Br \\ + \{(CH_2 = CCICH_3)_3NCH_3]Br$
	Amine	$C_{1}-C_{0}$ $(CII_{3})_{2}NCII_{2}CN$ $[(CII_{3})_{2}N]_{2}CH_{2}$ $(C_{2}I_{6})_{2}NCH_{2}CN$ $(n_{-}C_{3}H_{7})_{2}NCH_{3}CN$ $(n_{-}C_{3}H_{7})_{2}NCH_{3}$ $(n_{-}C_{3}H_{7})_{2}NCH_{2}CN$ $(n_{-}C_{3}H_{7})_{2}NCH_{2}CN$ $(C_{3}H_{5})_{2}NCH_{2}CO_{2}C_{2}H_{5}$ $(n_{-}C_{3}H_{7})_{3}N$	C ₁₀ -C ₁₅ [(CH ₃) ₂ CHCH ₂] ₂ NCH ₂ CN (n-C ₃ H ₇) ₂ NCH ₂ CO ₂ C ₂ H ₅ C ₆ H ₅ (CH ₂) ₂ N(CH ₃) ₂ C ₆ H ₅ (CH ₂) ₂ N(CH ₃) ₂ [(CH ₃) ₂ CHCH ₂] ₂ NCH(CH ₃) ₂ C ₆ H ₅ (CH ₂) ₂ N(C ₂ H ₅) C ₆ H ₅ (CH ₂) ₂ N(C ₂ H ₅) ₂	$[(CII_3)_2CHCH_2CH_2]_2NCH(CH_3)_2 \\ [(n-C_3H_7)_2N]_2CH_2 \\ C_6H_5(CH_2)_3N(C_3H_7-n)_2$	$C_{\tau}C_{\mathfrak{p}}$ $(CH_{2}=CClCH_{2})_{\mathfrak{p}}NCH_{3}$

OTAN	00221			
39 39 39 39 39 4 4	35 39 34	34	16 16 16 36	16 16
Mixed eyanamides + mixed bromides CH ₂ =CCICH ₂ N(CN)CH ₃ + BrCH=CHCH ₂ Br CH ₂ =CBrCH ₂ N(CN)CH ₃ + BrCH=CHCH ₂ Br CH ₂ =CBrCH ₂ N(CN)CH ₃ + CH ₂ =CBrCH ₂ Br Mixed cyanamides + mixed bromides CH ₂ =CBrCH ₂ N(CN)CH ₃ + CICH=CHCH ₂ Br CICH=CHCH ₂ N(CN)CH ₃ + CICH=CHCH ₂ Br (CH ₃) ₂ CH] ₂ NCN + CH ₂ =CHCH ₂ Br (C ₂ H ₅) ₂ NCN + amine hydrobromide ‡	(CH ₃) ₂ NCN + C ₆ H ₆ CH=CHCH ₂ Br ClCH=CHCH ₂ N(CN)CH ₃ + C ₆ H ₆ CH ₂ Br CH ₂ =CBrCH ₂ N(CN)CH ₃ + C ₆ H ₅ CH ₂ Br CH=CH CH=CH >CHN(CN)CH ₃ + amine hydrobromide ‡	$^{ m CH_2}$ —CH ₂ Inseparable mixture of two cyanamides $+$ amine hydrobromide \ddagger	$CH_3CH=CHCH_2N(CN)CH_3+p\text{-}CH_3C_6H_4CH_2Br$ $CH_2=CHCH_2N(CN)CH_3+C_6H_5CH=CHCH_2Br$ $CH_3CH=CHCH_2N(CN)CH_3+C_6H_5CH=CHCH_2Br$ $(C_6H_5CH_2)_2NCN+CH_2=CHCH_2Br$	$C_6H_5CH=CHCH_2N(CN)CH_3+p$ - $CH_3C_6H_4CH_2Br$ $C_6H_5CH=CHCH_2N(CN)CH_3+p$ - $C_6H_5C_6H_4CH_2Br$ at 100° along were not isolated.
CICH=CHCH ₂ N(CH ₃)CH ₂ CH=CHBr CH ₂ =CCICH ₂ N(CH ₃)CH ₂ CH=CHBr CH ₂ =CBrCH ₂ N(CH ₃)CH ₂ CH=CHBr CH ₂ =CCICH ₂ N(CH ₃)CH ₂ CBr=CH ₂ (CH ₂ =CCICH ₂ N(CH ₃)CH ₂ CBr=CH ₂ (CICH=CHCH ₂) ₂ NCH ₃ CICH=CHCH ₂ N[CH(CH ₃) ₂] ₂ CII =CH CHCHCCHC ₂ N[CH(CH ₃) ₂] ₂	ĠII₂—CĤ₂ CaHaCH=CHCH₂N(CH₃)₂ CICII==CHCH₂N(CH₃)CH₂CaHa CII₂==CBrCH₂N(CH₃)CH₂CaHa		$ \qquad \qquad \text{CH}_{2} \text{CH}_{3} \text{CH}_{3} \text{CH}_{3} \text{CH}_{3} \text{CH}_{4} \text{CH}_{2} \text{CH}_{3} \text{CH}_{3} \text{CH}_{2} \text{CH}_{4} \text{CH}_{2} \text{CH}_{4} \text{CH}_{3} \text{CH}_{2} \text{CH}_{3} \text{CH}_{2} \text{CH}_{4} \text{CH}_{4$	$C_{19}-C_{23}$ $p-CII_{3}C_{6}H_{4}CH_{2}N(CH_{3})CH_{2}CH=CHC_{6}H_{5}$ $p-CII_{3}C_{6}H_{4}CH_{2}N(CH_{3})CH_{2}CH=CHC_{6}H_{5}$ $p-C_{4}II_{5}C_{6}H_{4}CH_{2}N(CH_{3})CH_{2}CH=CHC_{6}H_{5}$ $p-C_{5}II_{5}C_{6}H_{4}CH=CHCH_{2}N(CN)CH_{3} + p-CI$ $p-C_{5}II_{5}C_{6}H_{4}CH=CHCH_{2}N(CN)CH_{3} + p-CI$ $p-C_{5}II_{5}C_{6}H_{4}CH=CHCH_{2}N(CN)CH_{3} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{2}N(CN)CH_{3} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{5}N(CN)CH_{3} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{5}N(CN)CH_{3} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{5}N(CN)CH_{5} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{5}N(CN)CH_{5} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{5}N(CN)CH_{5} + p-CI$ $p-C_{5}II_{5}C_{6}H_{5}CH=CHCH_{5}N(CN)CH_{5} + p-CI$ $p-C_{5}II_{5}CH=CHCH_{5}N(CN)CH_{5}N(CN)CH_{5} + p-CI$ $p-C_{5}II_{5}CH=CHCH_{5}N(CN)CH_{5$

ABLE III—Continued

				ORGANIC	REACTION	ONS
Refer-	ence		90 90 90	36 38 38	36	24 42 42 42 42 42 42 42 42 42 42 42 42 4
Acyclic Amines	Products	C. Derivatives of Benzylamine §	$(CH_3)_2NCN + {}_{\phi}\text{-}ClC_6H_4CH_2Br + \{(\phi\text{-}ClC_6H_4CH_2)_2N(CH_3)_2\}Br$	$(CH_3)_2NCN + p-1C_6H_4CH_2BT \ (C_3H_6)_2NCN + C_6H_5CH_2BT \ HC = CCH_2N(CN)CH_3 + C_6H_5CH_2BT \ (CH_3)_2NCN + o-CH_2 = CHC_6H_4CH_2BT \ (GH_3)_2NCN + o-CH_2 = C$	$CH_2 \longrightarrow CHCH_2N(CN)CH_3 + C_6H_6CH_2Br$ $(n-C_3H_7)_2NCN + C_6H_6CH_2Br$	Mixed eyanamides + mixed bromides p-ClC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-FC ₆ H ₄ CH ₂ Br Mixed cyanamides + mixed bromides Mixed cyanamides + mixed bromides m-ClC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-BC ₆ H ₄ CH ₂ Br m-BrC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-ClC ₆ H ₄ CH ₂ Br p-BrC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-ClC ₆ H ₄ CH ₂ Br p-BrC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-ClC ₆ H ₄ CH ₂ Br m-ClC ₆ H ₄ CH ₂ N(CN)CH ₃ + m-BrC ₆ H ₄ CH ₂ Br m-ClC ₆ H ₄ CH ₂ N(CN)CH ₃ + m-BrC ₆ H ₄ CH ₂ Br m-IC ₆ H ₄ CH ₂ N(CN)CH ₃ + m-BrC ₆ H ₄ CH ₂ Br m-IC ₆ H ₄ CH ₂ N(CN)CH ₃ + m-BrC ₆ H ₄ CH ₂ Br
	Amina	oming	$\mathrm{C_{9}\text{-}C_{13}}_{o\text{-}\mathrm{ClC}_{6}\mathrm{H}_{4}\mathrm{CH}_{2}\mathrm{N}(\mathrm{OH}_{3})_{2}}$	$\begin{array}{ll} p_{\text{-}1}\text{C}_{\text{6}}\text{H}_{\text{4}}\text{CH}_{\text{2}}\text{N}(\text{CH}_{\text{4}})_{\text{2}} \\ \text{C}_{\text{6}}\text{H}_{\text{6}}\text{CH}_{\text{3}}\text{N}(\text{C}_{\text{2}}\text{H}_{\text{6}})_{\text{2}} \\ \text{C}_{\text{6}}\text{H}_{\text{5}}\text{CH}_{\text{5}}\text{N}(\text{CH}_{\text{3}})\text{CH}_{\text{2}}\text{C} \\ \text{c}_{\text{-}C}\text{H}_{\text{2}}\text{CH}\text{C}_{\text{6}}\text{H}_{\text{4}}\text{CH}_{\text{2}}\text{N}(\text{CH}_{\text{3}})_{\text{2}} \end{array}$	$C_0H_2CH_2N(CH_3)CH_2CH_2$ $C_0H_3CH_2N(C_3H_7-n)_2$	C ₁₅ P-FC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₅ p-FC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₅ o-FC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₅ p-FC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₅ p-FC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ F-o p-BrC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m o-BrC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m p-BrC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m m-BrC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-p m-BrC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-p m-BrC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-p m-IC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m m-IC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m m-IC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m m-IC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl-m

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CANNOC	11214 1720	
# 9 9 9 9 9 9 9 9	555 48E588	ន្តភូន
o-IC ₆ II ₄ CH ₂ N(CN)CII ₃ + o-BrC ₆ II ₄ CII ₂ Br p-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + C ₆ II ₂ CII ₁ Br o-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + o-ClC ₆ II ₄ CII ₂ Br p-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + p-ClC ₆ II ₄ CII ₂ Br p-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + p-ClC ₆ II ₄ CII ₂ Br p-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + o-ClC ₆ II ₄ CII ₂ Br p-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + m-ClC ₆ II ₄ CII ₂ Br o-O ₂ NC ₆ II ₄ CII ₂ N(CN)CII ₃ + p-IC ₆ II ₄ CII ₂ Br Nixed cyanamides + mixed bromides Nixed cyanamides + mixed bromides	p-0 ₂ NC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-NCC ₆ H ₄ CH ₂ Br p-NCC ₆ H ₄ CH ₂ N(CN)CH ₃ + o-1C ₆ H ₄ CH ₂ Br p-NCC ₆ H ₄ CH ₂ N(CN)CH ₃ + C ₆ H ₅ CH ₂ Br + [(C ₆ H ₅ CH ₂ N(CN)CH ₃)CH ₂ C ₆ H ₄ CN-p]Br p-FC ₆ H ₄ CH ₂ N(CN)CH ₃ + p-CH ₃ C ₆ H ₄ CH ₂ Br C ₆ H ₅ CH ₄ CH ₅ N(CN)CH ₃ + p-CH ₃ C ₆ H ₄ CH ₂ Br C ₆ H ₅ CH ₂ N(CN)CH ₃ + p-CH ₃ C ₆ H ₄ CH ₂ Br o-CH ₃ C ₆ H ₄ CH ₂ N(CN)CH ₃ + p-CH ₃ C ₆ H ₄ CH ₂ Br C ₆ H ₆ CH ₂ N(CN)CH ₃ + m-CH ₃ C ₆ H ₄ CH ₂ Br	C ₆ H ₆ CH ₂ N(CN)C ₂ H ₅ + p-CH ₃ C ₆ H ₄ CH ₂ Br Mixed eyanamides + mixed bromides m-CH ₃ C ₆ H ₄ CH ₂ N(CN)CH ₃ + p-CH ₃ C ₆ H ₄ CH ₂ Br
		÷

 $m-O_2NC_6H_4CH_2N(CH_3)CH_2C_6H_4Cl-p$

 p_{-0} NC₆H₄CH₂N(CH₃)CH₂C₆H₄Cl-o

 $p_{-O_2\mathrm{NC}_6\mathrm{H_4CH_2N}}(\mathrm{CH_3})\mathrm{CH_2C}_6\mathrm{II_4Cl-p}$

o-O2NC6H4CH2N(CH3)CH2C6H4Cl-o

o-IC,H,CH2N(CH3)CH2C,H4Br-o

p-02NC6H4CH2N(CH3)CH2C6H6

p-0₂NC₆H₄CH₂N(CH₃)CH₂C₆H₄Cl-m

0-O2NC6H4CH2N(CH3)CH2C6H4I-0

 $p_{-}O_2NC_6H_4CH_2N(CH_3)CH_2C_6H_4CN-p$

o-IC,H,CH2N (CH3)CH2C,H,CN-p

p-NCC,H,CH2N(CH3)CH2C,H5

o-CH3C6H4CH2N(CH3)CH2C6H4Cl-p

p-CH₃C₆H₄CH₂N(CH₃)CH₂C₆H₆ p-CH₃C₆H₄CH₂N(CH₃)CH₂C₆H₆

m-CH₃C₆H₄CH₂N(CH₃)CH₂C₆H₅

 $C_{i,i}$

p-CH₃C₆H₄CH₂N(CH₃)CH₂C₆H₄F-p

p-CH₃OC₆H₄CH₂N(CH₃)CH₂C₆H₆

p-CH₃C₆H₄CH₂N(CH₃)CH₂C₆H₄CH₃-m

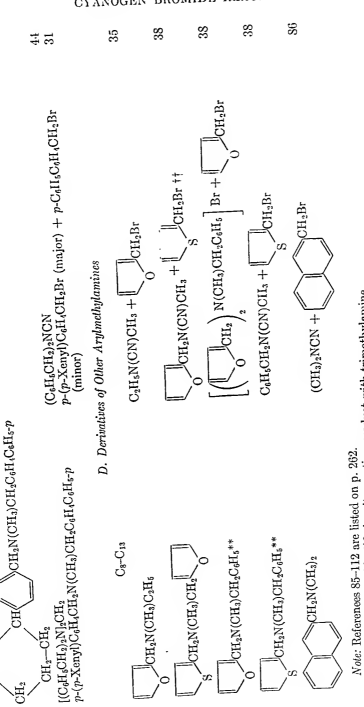
 $\begin{array}{ll} p\text{-}\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{N}(\mathrm{C}_2\mathrm{H}_5)\mathrm{CH}_2\mathrm{C}_6\mathrm{H}_5 \\ o\text{-}\mathrm{C}_2\mathrm{H}_5\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{N}(\mathrm{CH}_3)\mathrm{CH}_2\mathrm{C}_6\mathrm{H}_5 \end{array}$

[§] See also Section D, p. 239. || The products are not separable by distillation.

TABLE III-Continued

38					ORGA	NIC I	REACTIO	NS		
	Refer-	ence		23 23 23	24 40	940	404	16 31 31 40 40 1	31	23 16 16
	ACYCLIC AMINES	Products	C. Derivatives of Benzylamine—Continued	$ ho_{-\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{N}(\mathrm{CN})\mathrm{CH}_3+p_{-\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{Br}}$ $ ho_{-\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{N}(\mathrm{CN})\mathrm{CH}_3+m_{-\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{Br}}$ $p_{-\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{Br}}$	+ $(p$ -CH ₂ UC ₆ H ₄ CH ₂) ² M (CM ₂) ² CT ₂ C ₃ CT ₄ CH ₂	p-CH ₃ CONHC ₆ H ₄ CH ₂ N(CN)CH ₃ + C ₆ H ₆ CH ₃ Br + [(C ₆ H ₆ CH ₂) ₂ N(CH ₃)CH ₂ C ₆ H ₄ NHCOCH ₃ - p]Br	$p\text{-CH}_3\text{CONHG}_6\text{H}_4\text{CH}_2\text{N}(\text{CN})\text{CH}_3+p\text{-IC}_6\text{H}_4\text{CH}_2\text{DI}_3$ $m\text{-NCG}_6\text{H}_4\text{CH}_2\text{N}(\text{CN})\text{CH}_3+p\text{-NCG}_6\text{H}_4\text{CH}_2\text{BI}_3$ Mixed eyanamide + mixed bromide	p-CH ₃ C ₆ H ₄ CH ₂ N(CN)CH ₃ + p-C ₂ H ₆ C ₆ H ₄ CH ₂ Br Mixed cyanamides + mixed bromides Mixed cyanamides + mixed bromides p-CH ₃ CONHC ₆ H ₄ CH ₂ N(CN)CH ₃ p-CH ₃ CONHC ₆ H ₄ CH ₂ N(CN)CH ₃ CG ₆ H ₆ CH ₂ N ₂ N(CN)CH ₃ CG ₆ H ₆ CH ₂ N ₂ NCN + C ₆ H ₆ CH ₂ Br	CensCH2N(CN)CH3 + P-CensCentCusDi o-CeHeCeH4CH2Br ¶ + unidentified mixture of cyanamides	p-C ₆ H ₅ C ₆ H ₄ CH ₂ N(CN)CH ₃ + p -CH ₃ OC ₆ H ₄ CH ₂ Br p-CH ₃ C ₆ H ₄ CH ₂ N(CN)CH ₃ + p -C ₆ H ₅ C ₆ H ₄ CH ₂ Br p-C ₂ H ₅ C ₆ H ₄ CH ₂ N(CN)CH ₃ + p -C ₆ H ₅ C ₆ H ₄ CH ₂ Br
			Amine C. Deriva	C_{17} ($Cont^3$ d) o - $CH_3C_6H_4CH_2N(CH_3)CH_2C_6H_4CH_3-p$ o - $CH_3C_6H_4CH_2N(CH_3)CH_2C_6H_4CH_3-m$ o - $CH_3C_6H_4CH_3-p$	p-CH ₃ C ₆ H ₄ CH ₂ N(C ₂ H ₆)CH ₂ C ₆ H ₄ H ² - pp -CH ₃ CONHC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ Cl- p	$p\text{-}\mathrm{CH}_3\mathrm{CONHC}_6\mathrm{H}_4\mathrm{CH}_2\mathrm{N}(\mathrm{CH}_3)\mathrm{CH}_2\mathrm{C}_6\mathrm{H}_5$	p-CH ₂ CONHC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ I- op -NCC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ CN- mm -NCC ₆ H ₄ CH ₂ N(CH ₃)CH ₂ C ₆ H ₄ CN- o	$C_{18}-C_{21}$ $p-C_{2}H_{5}C_{6}H_{4}CH_{2}N(CH_{3})CH_{2}C_{6}H_{4}CH_{3}-p$ $p-CH_{2}OC_{6}H_{4}CH_{2}N(CH_{3})CH_{2}C_{6}H_{4}OC_{2}H_{5}-p$ $p-CH_{3}COHC_{6}H_{4}CH_{2}N(CH_{3})CH_{2}C_{6}H_{4}CH=CH_{2}-p$ $p-CH_{3}CONHC_{6}H_{4}CH_{2}N(CH_{3})CH_{2}C_{6}H_{4}NHCOCH_{2}-p$ $p-CH_{3}CONHC_{6}H_{4}CH_{2}N(CH_{3})CH_{2}C_{6}H_{4}NHCOCH_{2}-p$ $(C_{6}H_{5}CH_{2})_{3}N$	H ₂ N(CH ₃ H ₂ N(CH ₃	$p\text{-}\mathrm{CH}_s\mathrm{OC}_s\mathrm{H}_s\mathrm{CH}_z\mathrm{N}(\mathrm{CH}_s)\mathrm{CH}_z\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s\mathrm{C}_s\mathrm{H}_s\mathrm{C}_s$

31



** Though derivatives of benzylamine, these amines are listed in this section to emphasize the behavior of the lpha-furfuryl and †† The products were poorly characterized. α -thienyl groups.

This bromide was identified as its reaction product with trimethylamine.

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–Contin
-111
TABLE

		UKGAN	10 103	-		
Refer- ence	98	98	98	•	1, 2 45 45 41 1	43 1, 2 1
Acyclic Amines Products	D. Derivatives of Other Arylmethylamines—Continued CH ₂ Br	$(CH_3)_2NCN + (CH_3)_2NCN + $	+ C ₆ H ₆ CH ₂ Br	CH ₂ N(CN)CH ₃ +	E. Derpatences 9, 17.0 CoHoN(CN)CH3 + [CoHoN(CH3)3]Br m-ClCoH4N(CN)CH3 + [m-ClCoH4N(CH3)3]Br m-CH3CoH4N(CN)CH3 + [m-CH3CoH4N(CH3)3]Br m-CH3CoH4N(CN)CH3 + [m-CH3CoH4N(CH3)3]Br (ca. 75%) p-CH3CoH4N(CN)CH3 + [p-CH3CoH4N(CH3)3]Br (ca. 75%)	$C_{\rm eff, en}^{\rm cont}$ No reaction p -BrC _e H _e N(CH ₃)CH ₂ CN $C_{\rm eff, en}^{\rm cont}$ (CN)C ₂ H _e $C_{\rm eff, en}^{\rm cont}$ (CN)C ₂ H _e $C_{\rm eff, en}^{\rm cont}$ (CN)C ₂ H _e $C_{\rm eff, en}^{\rm cont}$
	Amine D .	CH ₂ N(CH ₃) ₂	CH ₂ N(CH ₃)CH ₂ C ₆ H ₅	CH ₂ N(CH ₃)CH ₂	Cg-C10 CgH5N(CH3)2 m-ClCgH4N(CH3)2 m-ClCgH4N(CH3)2	p-CH ₂ C ₂ L ₄ N(CH ₃)? C ₅ H ₅ N(CH ₃)C ₂ H ₅ p-BrC ₆ H ₄ N(CH ₃)CH ₂ CN C ₅ H ₅ N(CH ₃)CH ₂ CN ^{‡‡} C ₆ H ₅ N(C ₂ H ₅)?

8 88 84 38 38 43 43 3,H6N(CN)C3H7-i + amine hydrobromide $G_6H_5N(CN)G_3H_7-i+CH_2=CHCH_2Br$ Chunch + CH2=CHCH2Br O,H,N(CN)CH3+CH2-CHCH2Br C.H.S.N(CN)CH3+CH2=CHCH2Br (63-67%) $p_{-(i-C_3H_7)}C_6H_4N(CN)CH_3$ (37%) C,H,N(CN)CH3 + HC=CCH2Br $C_{\rm sH_5N}({\rm GN})C_{\rm sH_7-n}+n-C_{\rm sH_7Br}$ $C_6H_5N(CN)C_3H_7-i+n-C_3H_7Br$ C,H,N(CN)CH,CH-CH2 CH2-CH2 No definite products isolated CH_2 p-BrC6H4N(C2H5)CH2CN N(CN)CH3§§ No products isolated C, H, N(CN)C, H,-n C,H,N(CN)C,H-1 C,H,N(CN)C3H7-i No reaction C_{11} $-C_{13}$ C6H5N(C3H7-i)CH2CH=CH2 C,H,N(CH,)CH,CH-CH, CH2-CH2 CH2N(C2H4)CH2CH=CH2 C6H5N(CH3)CH2CH=CH2 p-CH3CeH4N(CH3)CH2CN CeH3)CH(CH3)CH(CH3)CN C,H,N(CH,)CH,C=CH p-(i-C3H7)C6H4N(CH3) C6H5N(C3H7-n)C3H7-1 O,H,N(C,H,)CH,CN ‡ C,H,N(CH3)CH2CH— 5,H,N(C,H,)C,H,-n 5,H,N(C,H,)C,H-i N(CH3)2 C6H3)C3H7-i 3,H5N(C3H7-n)2 $C_6H_5N(C_3H_7-i)_2$

 \ddag This reaction was carried out at 100°. No reaction occurs at room temperature. §§ The cthyl analog was obtained in 48% yield. Note: References 85-112 are listed on p. 262.

(C₆H₅)₂NCH₃

ACYCLIC AMINES

Reference

Products

38

 $J_{CH_2N}(CN)C_6H_6+CH_2=CHCH_2Br$

 $C_6H_6N(CN)C_4H_9-i+n-C_4H_9Br$

C14-C16

 $C_6H_6N(C_4H_9-n)C_4H_9-i$

Amine

E. Derivatives of Aromatic Amines—Continued

36 37 53

C,H,N(CN)CH3+ o-CH3C,H,CH2Br |||

C,H,N(CN)CH3 + C,H,CH2Br

CH2N(C6H8)CH2CH=CH2

C6H6(CH2)2N(CN)C6H6

CH2-CH2

33 36 36

Equal amounts of both cyanamides and bromides

 $C_6H_5N(GN)C_4H_9-n+CH_2-CHCH_2Br$

CH2-CH2

o-CH3C6H4CH2N(CH3)C6H5 $C_6H_5(CH_2)_2N(CH_3)C_6H_5$

C,H,N(CH3)CH2C,H5

n-C₄H₉N(C₆H₅)CH₂ĊH----ĊH₂

C6H6N(CN)CH2C6H6 + CH2=CHCH2Br

C,H,N(CN)C3H,-i + C,H,CH2Br

45,88

8

ĞeHeN(CN)CH2C=CH (60%) + n-CeH11C=CCH2Br ¶¶

[p-CH₃(CN)NC₆H₄]₂CH₂ (ca. 50%)

JN(CN)CH3 (45%)

CH3(CN)N

JN(CH3)2

 $HC = CCH_2N(C_6H_6)CH_2C = CC_6H_{11}$ -n

 $[p-(\mathrm{CH}_3)_2\mathrm{NC}_6\mathrm{H}_4]_2\mathrm{CH}_2$

 C_{17} - C_{19}

CeH,CH2N(CeH5)CH2CH=CH2

n-C₆H₁₁N(C₆H₆)C₆H₁₁-ii-C₃H₇N(C₆H₆)CH₂C₆H₆

N(CN)CH3

- - 45

 CH_{2}

45 46 46 45 N(CN)CH3 (ca. 40%) VN(CN)CH3 N(CN)CH CH_3 黑 CH3(CN)N No reaction CH3(CN)N \N(CH₃)₂ N(CH₃)2 $N(CH_3)_3$ CH_3 $\widetilde{\mathrm{CH}_3}$ E E (CH3)2N($(CH_3)_2N_3$

ĆH., [C6H6N(CN)]2(CH2)6 No products isolated

No reaction

No reaction

N(CH₃)2

EHI3

(CH3)2N

)CH?(

 CH_3

(C₆H₆)₃N [C₆H₅N(CH₃)]₂(CH₂)₅

(CH3)2N(

45

45

57

|||| The products were poorly characterized. ||¶ Appreciable cleavage in the other direction was observed. Note: References 85-112 are listed on p. 262. ĆH3

Refer-

ence

Products

E. Derivatives of Aromatic Amines—Continued

No reaction

 C_{12} (Cont'd)

CII,

Amine

No reaction

(CII.)

Acyclic Amines

45

An oil which on treatment with water yielded

 C_{20} – C_{24}

CH,

N(CIIs)2

56 68

 $\begin{array}{l} C_b H_s CH[C_b H_s N(CN) CH_{s^2} p]_2 \ (ca. \ 75\%) \\ C_b H_s C = CCH_s N(CN) C_b H_s(?) \end{array}$

. No reaction took place at the amino group. Note: References 85-112 are listed on p. 262.

[p-(CH3)3NC3H3]2HC3H5 C8H5CH=CHCH2N(C8H5)CH3C=CC6H5

45

45

25

Note: References 85-112 are listed on p. 262.

TABLE IV

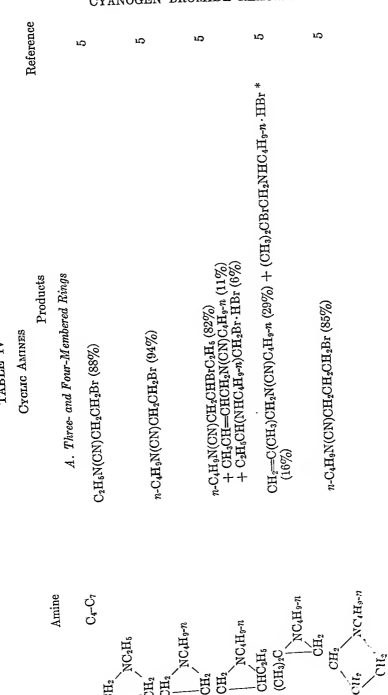


	TABLE IV—Continued	
	Cyclic Amnes Products	Reference
Amine	B. Five-Membered Rings	48
Cr-Cs	C ₂ II ₅ N(CN)(CH ₂),Br † (91%)	ł
C;11s	n-C ₃ 11,N(CN)(CH ₂),Br (93%)	48
Oallrn	n-C ₄ H ₉ N(CN)(CH ₂) ₄ Br (quant.)	5, 47
C ₄ II ₉ -n	(CH ₃) ₂ CHN(CN)CH(CH ₃)(CH ₃) ₃ Br (61%) + (CH ₃) ₂ CHN(CN)(CH ₂) ₃ CHBrCH ₃ (30%)	ಸು
CII(CII ₃) ₂	$_{n-\mathrm{C_6H_{11}N(CN)(CH_2)_4\mathrm{Br}}}(\mathrm{ca.~80\%})$	85
CsII11-n	;-C ₆ H ₁₁ N(CN)(CH ₂),Br	85
CsH11-i	$_{n}$ -C ₁ H ₅ N(CN)CH(CH ₂) ₃ Br (70%) + $_{n}$ -C ₄ H ₅ N(CN)(CH ₂) ₃ CHBrCH ₃ (26%) $_{CH_3}$	ಬ

n

20

CH₂CH₂Br (40%) (40%) CH₂Br ‡ CH₂Br ‡ CH₃Br ‡ CH₃Br ‡ CH₃Br ‡

C₉-C₁₁

 CH_3

CH₂=C(CH₂)₃N(CN)C₄H₉·n (41%) + amine hydrobromide (42%) CH₃

p-ClC₆H₄N(CN)(CH₂)₃CHBrCH₃ (ea. 50%) + p-ClC₆H₄N(CN)CH₂)₃CHBrCH₃ (ea. 10%)

 $NC_4H_{2^{-n}}$

(CH3);

NC,H9-n

Note: References 85-112 are listed on p. 262.

NC6H1CI-p

 CH_3

9 The primary bromide was isolated as its reaction product with diethylamine. The product was isolated as the piperidine derivative. The product was poorly characterized

20

23

43

Reference



CYCLIC AMINES

NCN (40%) + CII2 wat CHCH 2 Brt B. Fire-Membered Rings-Continued Products

NCHI CH-CH

C11. C15

Amina

Calisn(CN)(CII2)3CHBrCII3 (ca. 50%) + Calisn(CN)CH(CII2)3Br § (ca. 10%) p-CH3OC4HAN(CN)(CH2)3CHBrCH3 (cn. 45%) + p-CH3OC4HAN(CN)CH(CH2)3Br § (cn. 15%) ĊII3

NCN + 0-CH2-CHC6H4CH2Br

NCH, Call, CII-- CII-0

NCN + C6H5CH2Br1

+ 0-BrCH₂C₆H₄CH₂N(CN)CH₂C₆H₅

No definite products

C14-C21

NCII, Call, CIIrp

(CO,C,Us);

 6

53

[†] The products were poorly characterized. § The primary bromide was isolated as its reaction product with diethylamine.

54

63

TABLE IV-Continued

CYCLIC AMINES

Reference

5.4

C. Six- and Scren-Membered Rings-Continued

Amine

 C_{r} - C_{s}

CH,CH,CH,Br NON (

CH(CH3)CH2CH2Br NCN

Br

No definite products

79

က

58, 94

20

7, 57

44

NCN + CH2=CHCO2C2H5 + amine hydrobromide (43%) i-C₅H₁₁N(CN)(CH₂)₅Br

NCH2CH2CO2C2H6

NC,H11-i

 NC_4H_9-n

S

Br

No definite products

(CH₃)₂]

 $p\text{-BrC}_6\mathrm{H}_4\mathrm{N}(\mathrm{CN})(\mathrm{CH}_2)_5\mathrm{Br}$

C,H,N(CN)(CH2),Br

NCH(CN)C,Hon

CH3

Note: References 85-112 are listed on p. 262.

NC6H4Br-p

NC,H,

NCN + amine hydrobromide n-C₄H₉N(CN)(CH₂)₅Br

NCH(CH₃)2 C₉-C₁₁

52

Reference

83

CYCLIC AMINES

TABLE IV-Continued

C. Six- and Seven-Membered Rings-Continued

C11-C11

C:IIt

Amine

N (75%) CN

N(CN)C3Hrn CH,CH,CH,Br

N N

95

p-CH₃C₆H₄N(CN)(CH₂)₆Br

ŽĘ

 $C_3II_{7}n$

N. Cii

9

2

NCN + 0-C2H5C6H4CH2Br

NCH2C6H4C2H5-0

NCH(CN)CeH13-11

 C_{14} – C_{16}

No definite products

NCN + C,H,CH,Br

CH=CHCHBrC3H7-n

CH3.

N(CN)CH3

SS

 CH_3

62

$$\mathrm{NCH_2C_6H_4CH_3-}p$$

92

NCN (ca. 50%) $+ p - \text{CH}_3\text{C}_6\text{H}_4\text{CH}_2\text{B}_\Gamma$

Reference

3

33

49

35

29

TABLE IV-Continued

CYCLIC AMINES

Products

C. Six- and Seven-Membered Rings-Continued

C₆II₄OCII₅CII₂CII₂N(CN)(CII₂)₈Br (ca. 50%) + C₆H₆OCH₂CH₂Br

NCN + o-CII; =CIIC, H,CH,Br

NCN + & C2H6C6H4CH3Br

No definite products

S

29

(ca. 50%)

N(CN)CH3

CH3CH2CHBrC,H6

CII

CH,

Amire

Cit. Cit (Conf.d)

NCHICHERIOGARS

NCII, C,III, CIII --- CIII --

NCII2CoII1CII3-p

Remarks **

Reference 97, 98,

Reaction product of pyridine with cy-

$$\begin{array}{c|c} & & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

D. Pyridine-Type Amines

Amine $C_{i}^{-}C_{j}$

Products

ArNHCH=CHCH=CHCH=NAr·HBr

S

67, 68,

Water present in re-

action mixture

0

S

was treated with anogen bromide

an arylamine

S

** See pp. 218-219 for a description of the reactions involved in Table IVD. Note: References 85-112 are listed on p. 262.

TABLE IV-Continued

CYCLIC AMINES

Reference 2 2 65 75 Water present in re-Simultaneous reac-Simultaneous reaction with HCN tion with HCN action mixture Remarks H + amine hydrobromide H + amine hydrobromide D. Pyridinc-Type Amines-Continued Products NCN S Š ZZ OH S N Hypothetical Intermediate S Aming CrCis

Note: References 85-112 are listed on p. 262.

TABLE IV-Continued

Reference 68 89 89 Water present in reaction mixture Simultaneous reaction with HCN Remarks + amine hydrobromide 0 D. Pyridine-Type Amines-Continued Products CN Creuc Amnes Z C S Structure not given CN Hypothetical Intermediate S S H Cir Cis (Conf'd) Amine

TABLE V

		
	ALKALOIDS	
	Products	Reference
Amine		
C8-C15		99a
677	ОН	
HO CH 3	CH(CH ₃)CH ₂ CH ₂ Br	
VN ✓	X	
Retronecanol	ĈN	
CH2	^	76
~~~~	CH ₂	
	(CH ₂ ) ₄ Br	
VN ✓	CN CN	78
Lupinane		10
	NCN*	
NCH ₃	« [ /nex	
	<del></del>	
CO ₂ CH ₃	CO2CH8	
Anhydroecgonine methyl ester		
	O CH2CH2N(CN)CH3	61
,°~~		
H ₂ C NCH ₃	H ₂ C CH ₂ Br (ca. 55%)	
	+ quaternary salt (ca. 45%)	
Hydrohydrastinine	+ quaternary conviction	61
^	CH ₂ CH ₂ N(CN)CH ₃	
H ₂ C NCH ₃	CH2Dr (ca. 2070)	
O OCH ₃	O OCH3	
Hydrocotarnine	+ quaternary salt .	100, 101
	$C_{16}H_{24}N_3O_2Br$	
C ₁₅ H ₂₄ N ₂ O Lupanine	(	77
To the second se	$C_{17}H_{26}N_4B_{72} + C_{16}H_{25}N_3B_7$ (two isomers)	
C ₁₅ H ₂₆ N ₂ Sparteine		78
Sparteme	NCN (ca. 60%)	
C ⁶ H ² COO	C ₆ H ₆ COO (ca. 60%)	
Centros.	<b>&gt;</b>	
00 OH	$CO_2CH_3$	
CO ₂ CH ₃ Cocaine		
	1 O H. N.O. Br (7.5%)	102
C ₁₇ -C ₂₀	$C_{18}H_{20}N_3O_3Br$ (80%) + $C_{35}H_{40}N_5O_5Br$ (7.5%)	
C ₁₇ H ₂₀ N ₂ O ₃ 2,3-Diketonucidine	Br	74
CH _a N—CH ₂		
/	CH ₂ CH ₂ N(CN)CH ₃	
ĊH ₂	( )—( )	
	CH ₃ O OCH ₃	
( )—( )	O1130 D	
CH-O OCH;		
CH ³ O OCH ³		

Note: References 85-112 are listed on p. 262.

**Considerable ring cleavage occurred, and the yield of the product shown was small. See p. 223.

### TABLE V-Continued

#### ALKALOIDS

	Alkaloids	Ti-france
1. *	Products	Reference
Amine		74
C17-C20 (Con't)	CN CT	43
Tetrahydrothebaine	N—CH ₂	
	CH ₂	
	(_)	
	CH3O OCH3	
		103
$C_{19}H_{22}N_2O$	C ₂₀ H ₂₈ N ₃ O ₂ -2HBr	
Cinchonine	C20H22N 1OBr	104
C19H22N2O	C20H22N 2OB	
Cinchonidine	CN	105
CH ₃ COO N—CH ₂	CH ₂ COQ N—CH ₂	
CH ₃ COO CH ₂	CH ₂	
	$\prec \prec$	
	<b>⟨ ⟩</b> → <b>⟨</b> ⟩	
	CH30	
CH³O	$\alpha_{1}$ $\alpha$ $\alpha$	
Acetyldihydroöxycodeinone		104
$C_{20}H_{24}N_2O_2$	C22H24N4O2Br2	
Quinine	C22H24N4O2B12	103
C20H24N2O2	G22H24N4O2D12	
Quinidine C21		*05
$C_{21}H_{22}N_2O_2$	Addition product of undetermined composition	108
Strychnine		102
$C_{21}H_{24}N_2O$	$C_{22}H_{24}N_3OBr + (C_{22}H_{24}N_3OBr)_2$	102
Strychnidine		74
CH ₂ N	CH ₂ N(CN)CH ₃	
$\langle \rangle - CH_2 \rangle$	⟨	
CH³O OCOC	CH3 CH3O OCOCH3	
Acetyl-α-methylmorphim	ethine	
/==\	/=/	107
	$H_2N(CH_3)_2$ $CH_2N(CN)CH_3$	
$\langle \rangle - CH_2 \rangle$	$\langle \rangle \leftarrow CH_2 \rangle$	
CH³O OCO	CH, CH ₃ O OCOCH ₃	
Acetyl-8-methylmorphi	• 0	
Acetyl-6-methylmorphii	methine	

Acetyl-8-methylmorphimethine

Note: References 85-112 are listed on p. 262.

### TABLE V-Continued

#### ALKALOIDS

	ALKALOIDS	
	Products	Reference
Amine		
C ₂₁ (Con't)  CH ₃ N—CH ₂ CH ₂	CN N-CH ₂ CH ₂ (ca. 75%)	74
CH3COO OCOCH3	CH3COO OCOCH3	
Diacetylmorphine		400
C22-C25	RN(CN)CH2CH2OCH2CH2Br	108
RN		108
<u></u>	RN(CN)(CH ₂ ) ₆ Br	
RN	$C_{24}H_{26}N_3O_4Br + C_{47}H_{52}N_6O_8Br$	102, 106. 109, 110
C23H26N2O4	C24112611304-1	
Brucine	C24H28N3O4Br	109, 110
C23H28N2O4		73
Dihydrobrucine	$C_{24}H_{37}N_3 + C_{26}H_{46}N_2Br_2 + C_{24}H_{34}N_4 + conessine$	(0
C24H40N2		111
Conessine	C ₂ 4H ₃₇ N ₃ + C ₂ 6H ₄ 6N ₂ Br ₂ + C ₂ 4H ₃ 4N ₄ + isoconessine	111
C24H40N2	hydrobromide	*00
Isoconessine	CH₂Br	108
#		
RN	CH ₂ N(CN)R	
		112
C ₂₆ -C ₃₁	C27H26N2O4	
Acetylphenyldihydrothebaine $CH_3$ $N$ $C_6H_5COO$ $OCOC_6H_5$	$CH_2CH_2N(CN)CH_3$ $C_6H_5COO$ $OCOC_6H_5$	75
Dibenzoylapomorphine		
Note: References 85-112 are listed on p. 2	262.	
Mote: References 55-112 are CH ₂ C	*Ho-	
CH ₂ C	)12/2	

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#### CHAPTER 5

### HYDROGENOLYSIS OF BENZYL GROUPS ATTACHED TO OXYGEN, NITROGEN, OR SULFUR

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and

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Philadelphia, Pa.

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phate 263	•
200	

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#### INTRODUCTION

The benzyl group and a variety of substituted benzyl groups attached to an oxygen atom as in alcohols, ethers, acetals, or esters; to an amino nitrogen atom; or to a sulfur atom in thio ethers may be removed as toluene, or the correspondingly substituted toluene, by hydrogenolysis.

$$ArCH_2OH + H_2 \rightarrow ArCH_3 + H_2O$$
  
 $ArCH_2OR + H_2 \rightarrow ArCH_3 + ROH$   
 $ArCH(OR)_2 + 2H_2 \rightarrow ArCH_3 + 2ROH$ 

application is the synthesis of phenylacetic acid from the acetate of mandelic acid.

$$C_6H_5CHCO_2H + H_2 \rightarrow C_6H_6CH_2CO_2H + CH_3CO_2H$$
  
 $OCOCH_3$ 

It is the purpose of this chapter to give illustrations of both types of debenzylations so that the usefulness of the reactions may be better appreciated and their applications extended. Since most descriptions of debenzylations in the literature are subordinated to other aspects of the studies in which they are reported, it is certain that not all of the examples of the reaction have been found and discussed in the text or listed in the tables.

#### SCOPE AND LIMITATIONS

Substituents may be present in the methylene side chain or in the nucleus of the benzyl group. The effects of the various substituents, in either the methylene or the phenyl group, are best considered under the various types of debenzylations as discussed in the following subsections: removal of the benzyl attached to oxygen, to nitrogen, or to sulfur.

The role of the benzyl group may also be taken by the benzhydryl ¹⁴ or the triplienylmethyl ¹⁵ group.

Hydrogenolysis may be accomplished by either chemical or catalytic means. Palladium seems to be the favored catalyst, but platinum, nickel, and copper chromium oxide have also been used successfully. No study of their relative merits has appeared. Chemical debenzylations have been effected by Rancy nickel alloy, sodium amalgam, sodium in liquid ammonia, and lithium aluminum hydride.

¹¹ Suter and Ruddy, J. Am. Chem. Soc., 66, 747 (1944).

¹¹ Michael, Ber., 65, 262 (1932).

# Cleavage of the Benzyl-Oxygen Bond

Alcohols, aldehydes, and ketones (Tables I, II, III, and IV). Benzyl alcohol is rapidly and quantitatively reduced to toluene. Nuclear-substituted benzyl alcohols behave similarly. p-Methoxybenzyl alcohol in ethanolic solution on reduction with palladium on charcoal forms p-methylanisole, 16 and salicin reduced with colloidal platinum, 17 platinum black, or palladium black  18  furnishes o-tolylglucoside.

$$p$$
-CH₃OC₆H₄CH₂OH + H₂  $\rightarrow p$ -CH₃OC₆H₄CH₃ + H₂O

Cinnamyl alcohol, a vinylog of benzyl alcohol, is reduced by hydrogen and palladium-carbon catalyst to a mixture of n-propylbenzene and 3-phenyl-1-propanol.¹⁶ It is probable, by analogy with information on nuclear hydrogenation,19 that these products result from competing and not from successive reactions: hydrogenation of the ethylenic bond to furnish the alcohol and "decinnamylation" by hydrogenolysis, followed by reduction of the double bond to furnish propylbenzene.

Benzyl alcohols substituted in the  $\alpha$  position likewise undergo hydrogenolysis. 1-Phenyl-1-propanol is reduced to propylbenzene, 20 1-phenyl-1-ethanol forms ethylbenzene, 21 1-phenylethane-1,2-diol yields phenethyl alcohol, and diphenylcarbinol is converted to diphenylmethane. 16

Since aldehydes of the general formula ArCHO may be reduced to the corresponding benzyl alcohols, ArCH2OH, and ketones of general structure ArCOR form  $\alpha$ -substituted benzyl alcohols, ArCHROH, it is to be expected that many aldehydes and ketones may be reduced directly to the corresponding toluenes or alkylbenzenes without the isolation of the intermediate alcohol. This expectation is realized in practice. 16, 20, 22, 23 Many aldehydes and ketones have been reduced at room temperature and low pressures to the corresponding hydrocarbons with hydrogen and palladium-carbon or copper chromium oxide cata-

¹⁶ Baltzly and Buck, J. Am. Chem. Soc., 65, 1984 (1943).

¹⁷ Kariyone and Kondo, J. Pharm. Soc. Japan, 48, 684 (1928) [C. A., 23, 393 (1929)].

¹⁸ Richtmyer, J. Am. Chem. Soc., 56, 1633 (1934).

¹⁹ Van Duzee and Adkins, J. Am. Chem. Soc., 57, 147 (1935).

²⁰ Hartung and Crossley, J. Am. Chem. Soc., 56, 158 (1934).

²¹ Kindler, Scharfe, and Henrich, Ann., 565, 51 (1949).

²³ Hartung and Smith, J. Elisha Milchell Society, 66, 171 (1950) [C. A., 47, 2716 (1953)]. ²² Hartung and co-workers, unpublished results.

lysts (Table III). Similar results may be accomplished by using Raney nickel-aluminum alloy and alkali.24

If the aryl alkyl ketone contains a phenolic hydroxyl in the ortho position, reduction to the hydrocarbon derivative does not take place. o-Hydroxypropiophenone is not reduced by palladium-carbon catalyst, and the 4-acylresorcinols are not reduced to the corresponding alkylresorcinols by either palladium or Raney nickel.22 For such reductions the Clemmensen ²⁵ or the Wolff-Kishner ²⁶ reactions must be used. Also complete substitution in the  $\alpha$  position of the aryl alkyl ketone inhibits hydrogenolysis. Pivalophenone, C6H5COC(CH3)3, is smoothly and quantitatively reduced to the carbinol but not to the hydrocarbon.16 The same behavior may be expected from other aryl t-alkyl ketones.

The hydrochlorides of aryl α-aminoalkyl ketones, ArCOCHRNH₃Cl, are reduced only to the amino alcohol when palladium catalyst is employed; however, if the amino ketone or the amino alcohol is hydrogenated in acetic acid at 80-90° with palladium on barium sulfate in the presence of perchloric acid, excellent yields of the desoxy compound are obtained.27 It is suggested that in the presence of perchloric acid the reduction proceeds through the acetic acid ester of the amino alcohol.

$$\begin{array}{c} \text{Arcochrnh}_3\text{Cl} \\ \text{H}_2 \downarrow \text{Pd} \end{array} \longrightarrow \begin{array}{c} \text{Pd, H}_2, \text{HClO}_4 \\ \text{CH}_3\text{CO}_2\text{H} \end{array} \longrightarrow \begin{array}{c} \text{Arch}_2\text{Chrnh}_3\text{Cl} \end{array}$$
 Archohchrnh $_3\text{Cl}$ 

An extension of the development described in the preceding paragraph is the reduction in one step, by means of palladium catalyst in acetic acid-perchloric acid solution, of  $\alpha$ -oximino ketones to the corresponding amines.27 The reduction of benzaldehyde cyanohydrin to phenethyl-

amine does not require the presence of acetic or perchloric acid but proceeds in ethanolic hydrogen chloride solution.28

The reduction of esters of aromatic acids to the corresponding hydrocarbons by means of copper chromium oxide 29 occurs by virtue of the

$$ArCO_2C_2H_5 + 3H_2 \rightarrow ArCH_3 + C_2H_5OH + H_2O$$

fact that these esters are first reduced to the aromatic alcohols, and the alcohol then undergoes hydrogenolysis. Ethyl benzoate, for example, reduced with copper chromium oxide in methanolic solution at 300 atm.

²⁴ Papa, Schwenk, and Whitman, J. Org. Chem., 7, 587 (1942).

Martin, in Adams, Organic Reactions, Vol. I, p. 155, John Wiley & Sons, 1942.

Todd, in Adams, Organic Reactions, Vol. IV, p. 378, John Wiley & Sons, 1948.

²⁷ Rosenmund and Karg, Ber., 75, 1850 (1942). 3 Hartung, J. Am. Chem. Soc., 50, 3370 (1928).

³ Laxier, U. S. pat. 2,079,414 [C. A., 31, 4340 (1937)].

and 125-175° is converted to benzyl alcohol.³⁰ If the temperature is increased to 200-250°, the products of the reaction are toluene, ethanol, and water.³¹

The ability of lithium aluminum hydride to effect hydrogenolysis of benzyl alcohols bearing an amino substituent in the *ortho* or *para* position is a recent discovery.³² Since this reducing agent converts esters, aldehydes, or ketones to carbinols,^{32a} it is seen that appropriately substituted intermediates may be converted directly to the corresponding toluidines. Illustrative of this reaction are the conversion of methyl anthranilate to o-toluidine (39%), o-aminobenzyl alcohol to o-toluidine (53%), p-aminobenzoic acid to p-toluidine (47%), p-dimethylaminobenzaldehyde to N,N-dimethyl-p-toluidine (78%), and p-aminobenzophenone to p-aminodiphenylmethane (57%).

Ethers (Table V). Hydrogenolysis of benzyl ethers proceeds smoothly, and the yields of products are generally good. Nickel or platinum catalysts may be used, but palladium is preferred if hydrogenation of the nucleus is to be avoided.

Benzyl alkyl ethers are quantitatively reduced to toluene and the corresponding alcohol by palladium ¹² or by Raney nickel. ¹⁹ Benzyl phenyl ether is converted into toluene and phenol when palladium-charcoal catalyst is used; ¹¹ but with Raney nickel as catalyst at 100° and 150–200 atm. toluene and both phenol and cyclohexanol are formed. ¹⁹

The hydrogenolyses described in the preceding section, where the benzyl group is retained in the product desired, have their parallel in certain oxygen heterocycles containing an  $\alpha$ -phenyl substituent, for example, the conversion of 2-phenyltetrahydropyran into 5-phenyl-1-pentanol and of phenyldioxane into phenethyl  $\beta$ -hydroxyethyl ether.²³

$$\begin{array}{c|c} CH_2 \\ H_2C & CH_2 \\ & \mid & \mid \\ H_2C & CHC_6H_5 \end{array} \longrightarrow C_6H_5(CH_2)_5OH$$

$$\begin{array}{c|c} O \\ O \\ \hline \\ H_2C & CH_2 \\ & \mid & \mid \\ H_2C & CH_2 \\ & \mid & \mid \\ H_2C & CHC_6H_5 \end{array} \longrightarrow C_6H_5CH_2CH_2OCH_2CH_2OH$$

³⁰ Mozingo and Folkers, J. Am. Chem. Soc., 70, 229 (1948).

³¹ Adkins, Reactions of Hydrogen, pp. 97-104, University of Wisconsin Press, 1937.

²² Conover and Tarbell, J. Am. Chem. Soc., 72, 3586 (1950).

²² Brown, in Adams, Organic Reactions, Vol. VI, p. 469, John Wiley & Sons, 1951.

²² Baker, Cornell, and Cron, J. Am. Chem. Soc., 70, 1490 (1948).

The principal application of the hydrogenolysis of benzyl ethers is in removing a benzyl group introduced earlier in order to protect a hydroxyl group during a series of reactions. For example, 1-(3-methoxy-4-benzyloxyphenyl)-2-acetaminopropanol (I) may be cyclized to the isoquinoline derivative II and the benzyl group then removed by hydrogenolysis to liberate the hydroxyl group in the 7 position of the isoquinoline III.³⁴ 6,7-Dihydroxy-1-(3',4'-methylenedioxybenzyl)isoquinoline (IV) may be prepared in an analogous manner.³⁵

$$\begin{array}{c} \text{CH}_3\text{O} \\ \text{C}_6\text{H}_5\text{CH}_2\text{O} \\ \text{C}_6\text{H}_5\text{C}_6\text{H}_2\text{O} \\ \text{C}_6\text{H}_3\text{C}_6\text{H}_2\text{O} \\ \text{C}_6\text{H}_3\text{C}_6\text{H}_3\text{C}_6\text{H}_3\text{C}_6\text{H}_3\text{C}_6\text{C}_6\text{H}_3\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C}_6\text{C$$

For the preparation of 3-(7-hydroxy-n-heptyl)veratrole (V) the Grignard reagent from 6-benzyloxy-1-bromohexane was allowed to react with 2,3-dimethoxybenzaldehyde to form a carbinol, which was dehydrated; reduction of the unsaturated intermediate in acetic acid solu-

tion with palladium black saturated the double bond and simultaneously removed the benzyl group.36

removed the benzy's group 
$$\begin{array}{c} OCH_3 \\ OCH_$$

The benzyloximino compounds are also useful in masking oximes because of the ease with which the protecting benzyl group may be removed by hydrogenolysis.  $\alpha$ -Oximino acids cannot be converted into their corresponding acid chlorides, but the O-ethers, the alkyloximino acids, are conveniently available and can be converted in good yields into the corresponding acid chlorides by the usual methods.37 The  $\alpha$ -benzyloximino acid chlorides react with  $\alpha$ -amino acids to form amides (VI) which may be reduced to dipeptides; 38 and the acid chloride will react with a dipeptide to form an attractive intermediate (VII) for the synthesis of a tripeptide.39

RCCOCl 
$$\xrightarrow{\text{NH}_2\text{CH}_2\text{CONHCH}_2\text{CO}_2\text{H}} \\ \text{RCCONHCH}_2\text{CONHCH}_2\text{CO}_2\text{H} \\ \text{NOCH}_2\text{C}_6\text{H}_5 \\ \text{VII} \\ \text{NOCH}_2\text{C}_6\text{H}_5 \\ \text{VII} \\ \text{RCCONHCHR'CO}_2\text{H} \\ \text{RCCONHCHR'CO}_2\text{H} \\ \text{RCH(NH}_2\text{)CONHCH}_2\text{CONHCH}_2\text{CO}_2\text{H} \\ \text{NOCH}_2\text{C}_6\text{H}_6 \\ \text{VI} \\ \text{H}_2\downarrow\text{Pd} \\ \text{RCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CO}_2\text{H} \\ \text{RCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CO}_2\text{H} \\ \text{NOCH}_2\text{C}_6\text{H}_6 \\ \text{VI} \\ \text{H}_2\downarrow\text{Pd} \\ \text{RCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CONHCH}_2\text{CO}_2\text{H} \\ \text{RCH}_2\text{CONHCH}$$

RCH(NH₂)CONHCHR'CO₂H

Acetals (Table VI). Hydrogenolysis of acetals of benzaldehyde furnishes toluene and the alcohol from which the acetal was formed. 10,40

and the abolics 
$$C_6H_5CH(OR)_2 \xrightarrow{H_2} C_6H_5CH_3 + 2ROH$$

²⁵ Wasserman and Dawson, J. Org. Chem., 8, 73 (1943).

³⁷ Waters and Hartung, J. Org. Chem., 12, 469 (1947).

³⁸ Weaver and Hartung, J. Org. Chem., 15, 741 (1950).

³⁹ Kramer, Hartung, and Hager, Chicago Meeting, American Chemical Society, September 1950.

⁴⁰ Sigmund, Monatsh., 53-54, 607 (1929).

The reaction is useful for the preparation of otherwise inaccessible esters of certain polyhydroxy compounds, for example, the  $\beta$ -monoglycerides. Glycerol and benzaldehyde form the 1,3-diacetal, leaving the secondary alcoholic group available for esterification; hydrogenolysis of the benzal group affords toluene and the  $\beta$ -glyceride. The benzaldehyde acetals of

sugars undergo similar hydrogenolyses. Benzal- $\alpha$ -methylglucoside with hydrogen in the presence of platinum sponge forms toluene and  $\alpha$ -methylglucoside.⁴²

Benzaldehyde diacetate has been reduced to toluene and acetic acid.⁷ No practical applications of this type of hydrogenolysis have been reported.

Esters (Tables VII and VIII). Esters of benzyl alcohol are reduced practically quantitatively to toluene and the acid from which the ester is formed.^{7,9} The reduction of the acetates of mandelie acid and its nuclear-substituted derivatives to the corresponding arylacetic acids, by means of palladium on barium sulfate and hydrogen, illustrates the type of hydrogenolysis in which the product of interest retains the benzyl group.⁴²

$$\begin{array}{c} \text{ArCHCO}_2\text{H} \rightarrow \text{ArCH}_2\text{CO}_2\text{H} + \text{CH}_3\text{CO}_2\text{H} \\ \downarrow \\ \text{OCOCH}_3 \end{array}$$

Hydrogenolyses of benzyl esters have also found important use in syntheses in which benzyl groups are employed to protect earboxyl groups and hence are not retained in the final products. Alkaline hydrolysis of an acylated malonic ester such as  $RCOCR'(CO_2C_2H_5)_2$  does

Bergmann and Carter, Z. physiol. Chem., 191, 211 (1930).
 Freudenberg, Toepfer, and Anderson, Ber., 61, 1750 (1928).

a Rosenmund and Schindler. Arch. Pharm., 256, 281 (1928).

not lead to the corresponding malonic acid for the acyl group is hydrolyzed more rapidly than the ester groups. 43a The benzyl esters, however, submit smoothly to hydrogenolysis with palladium-charcoal; decarboxylation of the malonic acid affords the ketone.44 This method has

been employed for the synthesis of compounds such as 3-tridecanonoic acid, 8-heptadecanone, 14-ethyl-13-octadecanonoic acid, 11-eicosanon-1-ol, 1-phenyl-2-pentanon-1-ol, and 3-m-methoxybenzoylpropionic acid.

A most attractive use of the debenzylation of esters by hydrogenolysis is the carbobenzyloxy method, developed by Bergmann and Zervas, 45, 46 for the synthesis of the peptide linkage. Carbobenzyloxy chloride,  $C_6H_5CH_2OCOCl$ , reacts with an amino acid to form a benzyl carbamate, C₆H₅CH₂OCONHCHRCO₂H; the free carboxyl group in this product may be converted into an acid chloride function, which by reaction with another molecule of amino acid yields the intermediate for a dipeptide. Hydrogenolysis forms toluene and a carbamic acid which

 $\text{C}_6\text{H}_6\text{CH}_2\text{OCONHCHRCO}_2\text{H} \rightarrow \text{C}_6\text{H}_6\text{CH}_2\text{OCONHCHRCOCl} \xrightarrow{\text{NH}_2\text{CHR}'\text{CO}_2\text{H}}$ 

 $C_6H_6CH_2OCONHCHRCONHCHR'CO_2H \xrightarrow{H_2}$ 

NH₂CHRCONHCHR'CO₂H + C₆H₅CH₃ + CO₂

spontaneously loses carbon dioxide, thus liberating the amino group which was protected during formation of the peptide linkage. The hydrogenolysis is effected by palladium black and hydrogen, and the yields are generally good. The free carboxyl group of the dipeptide derivative may, via its acid chloride, be coupled with a third amino acid, and so on, debenzylating only at the end of the synthesis.⁴⁷ An indication of the extent to which this reaction has been applied is shown in Table VIII.

The p-bromobenzyl carbamates, prepared from amino acids and p-bromocarbobenzyloxy chloride, have higher melting points and crystallize better than the corresponding benzyl carbamates. The p-bromo

The acid hydrolysis and decarboxylation of the acylated malonic ester C2H6O2-CCH₂CH₂COCH(CO₂C₂H₆)₂ to the acid HO₂CCH₂CH₂COCH₂CO₂H has been carried out by Eisner, Elvidge, and Linstead, J. Chem. Soc., 1950, 2223.

⁴⁴ Bowman, J. Chem. Soc., 1950, 325.

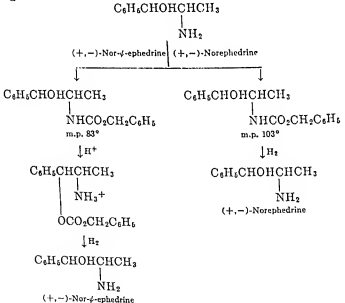
Bergmann and Zervas, Ber., 65, 1192 (1932).

⁶ Bergmann and Zervas, Ber., 65, 1201 (1932).

⁴⁷ Barkdoll and Ross, J. Am. Chem. Soc., 66, 951 (1944).

derivatives undergo hydrogenolysis in the same manner as do the unhalogenated carbamates.^{47a}

Because of the mild conditions under which benzyl carbamates respond to hydrogenolysis, certain derivatives lend themselves well for the recovery of pure isomers from a mixture of diastereoisomeric carbamates, thus avoiding the risk of Walden inversion or other chemical reactions which may accompany chemical deacylations. This is illustrated by the separation of the two racemic forms of norephedrine by way of their carbobenzyloxy derivatives. (+, -)-Nor- $\psi$ -ephedrine forms a urethane in which the amide group migrates quantitatively from the nitrogen to the oxygen atom, thus permitting easy separation of N-carbobenzyloxy-(+, -)-norephedrine from O-carbobenzyloxy-(+, -)-nor- $\psi$ -ephedrinc. Hydrogenolysis of each derivative regenerates the corresponding racemate.



The carbobenzyloxy method promises to be useful for the synthesis of aminoalkylmalonic acids, NH₂CR(CO₂H)₂. Aminomalonic ester, first converted into its carbobenzyloxy derivative, can be alkylated; the ethyl ester groups may be removed by milder hydrolysis than the benzyl ester, thus forming a carbobenzyloxyaminoalkylmalonic acid (VIII); the

 ⁴⁷a Channing, Turner, and Young, Nature, 167, 487 (1951).
 48 Fodor and Kiss. Nature, 163, 287 (1949).

mild conditions of the hydrogenolytic reaction permit reduction of the malonic acid or its salt.49

$$\begin{array}{c} C_{6}H_{5}CH_{2}OCONHCH(CO_{2}C_{2}H_{5})_{2} \xrightarrow{Alkylation} \\ \\ R \\ \downarrow \\ C_{6}H_{5}CH_{2}OCONHC(CO_{2}C_{2}H_{5})_{2} \xrightarrow{Mild \ alkaline \ hydrolysis}} C_{5}H_{5}CH_{2}OCONHC(CO_{2}H)_{2} \xrightarrow{H_{2}} \\ \\ R \\ \downarrow \\ VIII \\ R \\ NH_{2}C(CO_{2}H)_{2} + C_{6}H_{5}CH_{3} + CO_{2} \end{array}$$

The carbobenzyloxy group can also be removed by chemical means. Carbobenzyloxy-β-alanine, treated with sodium in liquid ammonia, is converted into  $\beta$ -alanine and 1,2-diphenylethane. 50

$$2C_6H_5CH_2OCONHCH_2CH_2CO_2H \rightarrow$$

$$2NH_2CH_2CH_2CO_2H + C_5H_5CH_2CH_2C_6H_5 + 2CO_2$$

The benzyl esters of phosphoric acid are employed to admirable advantage in the synthesis of phosphorylated amines and alcohols. 60x-6 #1 The general equations may be summarized as follows.

general equations may be seen as general equations may be seen as 
$$(C_6H_5CH_2O)_2POH \xrightarrow{Cl_2} (C_6H_5CH_2O)_2POC!$$

1.  $C_6H_5CH_2OH \xrightarrow{PCl_3} (C_6H_5CH_2O)_2POH \xrightarrow{H_2} (C_6H_5CH_2O)_2POC!$ 

1. 
$$C_6H_5CH_2OH \longrightarrow (C_6H_5CH_2O)_2PONR_2 \xrightarrow{H_2} R_2NPO_3H_2$$
  
2.  $(C_6H_5CH_2O)_2POCI \xrightarrow{R_2NH} (C_6H_5CH_2O)_2PONR_2 \xrightarrow{H_2} R_2NPO_3H_2$ 

2. 
$$(C_6H_6CH_2O)_2POCI \xrightarrow{HOR} (C_5H_6CH_2O)_2POOR \xrightarrow{H_2} ROPO_3H_2$$
  
3.  $(C_6H_6CH_2O)_2POCI \xrightarrow{HOR} (C_5H_6CH_2O)_2POOR \xrightarrow{H_2} ROPO_3H_2$ 

The mild conditions under which hydrogenolysis is effected make possible the synthesis of phosphorylated products of biological significance, which heretofore could be obtained with difficulty or by ambiguous procedures.

# Cleavage of Benzyl-Nitrogen Bonds

Amines (Tables IX-XV). Benzylamine, unlike benzyl alcohol, does not readily undergo hydrogenolysis. With palladium oxide 11 or with palladium-charcoal 22 no reduction was observed, and with nickel and

Beaujon, M.S. thesis, University of North Carolina, 1950.

⁵⁰ Sifford and du Vigneaud, J. Biol. Chem., 108, 753 (1935). 553 Atherton, Openshaw, and Todd, J. Chem. Soc., 1945, 382, 660.

¹²³ Atherton and Todd, J. Chem. Soc., 1947, 674.

^{1:}c Atherton, Howard, and Todd, J. Chem. Soc., 1948, 1106. and Baddiley, Clark, Michalski, and Todd, J. Chem. Soc., 1949, 815.

^{5:4} Michelson and Todd, J. Chem. Soc., 1949, 2476, 2487.

^{*} Michelson and Todd, J. Chem. Deen syl esters of phosphoric acid was described by Zervas, Naturwissenschaften, 27, 317 (1939).

n Birkofer, Ber., 75, 429 (1912).

hydrogen at high temperatures the hydrogenolysis was slight.^{1,2} Secondary amines containing one benzyl and one alkyl group also appear not to undergo hydrogenolysis; ^{15,51,52} in fact, one general method for preparing benzylamines of this type is the catalytic hydrogenation of the intermediate Schiff bases.^{52a} The following secondary amines were also found to be resistant to debenzylation: C₆H₅CH₂NH(CH₂)₃COCH₃ and C₆H₅CH₂NH(CH₂)₃CH(CH₃)NH₂. The latter, however, after conversion to the dimethylamino derivative with formaldehyde and formic acid did cleave at the benzyl-nitrogen bond to form NH₂(CH₂)₃CH-(CH₃)N(CH₃)₂.⁵³ The heterocyclic compounds IX and X were stable as hydrochlorides, but the free base IX underwent hydrogenolysis.^{53,54}

Certain secondary amines containing a benzyl group and an alkyl group which itself carries a non-hydrocarbon substitutent do undergo debenzylation to yield the corresponding primary amine; e.g.,

 $C_5H_5CH_2NHCH_2CH_2CO_2H \ ^{57} \qquad C_6H_6CH_2NHCH(CH_3)CH_2OH \ ^{58} \\ C_5H_5CH_2NHCHCO_2H \ ^{57} \qquad CH_3(CH_2)_3CH(NHCH_2C_5H_5)CH_2OH \ ^{58} \\ C_6H_5CH_2NHCHCO_2H \\ \\$ 

Secondary amines containing an aryl and a benzyl group are readily reduced to toluene and the primary aromatic amines. 7, 13, 51

Dibenzylamine is resistant to hydrogenolysis; it can in fact be prepared in 97% yield by the reduction of tribenzylamine with palladium oxide. However, dibenzylamines in which one benzyl group is substituted in the aromatic nucleus are amenable to hydrogenolysis, the unsubstituted benzyl group being removed. By means of competitive debenzylation studies (Table XII) of a series of 4,4′-disubstituted

⁵² Buck and Baltzly, J. Am. Chem. Soc., 63, 1964 (1941).

Emerson, in Adams, Organic Reactions, Vol. IV, p. 174, John Wiley & Sons, 1948.

Eisleb and Ehrhart, Ger. pat. 550,762 (Chem. Zentr., 1932 II, 615).
 Burger and Deinet, J. Am. Chem. Soc., 67, 566 (1945).

Mattocks and Hartung, J. Am. Chem. Soc., 68, 2108 (1946).

⁵⁶ Chemische Fabrik vorm. Sandoz, Fr. pat. 844,225 [C. A., 34, 7296 (1940)]; Peyer, U. S. pat. 2,243,977 [C. A., 35, 5508 (1941)].

⁵⁷ Wenner, U. S. pat. 2,389,099 [C. A., 40, 1539 (1946)].

is Niemann and Redemann, J. Am. Chem. Soc., 68, 1932 (1946).

$$\begin{array}{c}
\text{OH} & \xrightarrow{\text{CH}_2\text{O}} & \xrightarrow{\text{CH}_3\text{OH}} & \text{CH}_3\text{OH} \\
\text{OH} & \xrightarrow{\text{CH}_3\text{PNH}} & \text{CH}_3\text{PNH}_2\text{COH} & \text{OH} \\
& & \text{OH} & \text{OH} \\
& & \text{OH} & \text{OH}
\end{array}$$

Quaternary Ammonium Compounds (Table XV). Little attention has been given to the hydrogenolysis of quaternary benzylammonium compounds. Tribenzylmethylammonium hydroxide reduced with palladium oxide furnishes toluene and benzylmethylamine.⁵¹ Benzylphenyldimethylammonium chloride under similar conditions forms cyclohexyldimethylamine,⁵¹ an unusual instance of the reduction of the benzene nucleus with a palladium catalyst.

Chemical hydrogenolysis of quaternary ammonium compounds has received more study, which chronologically preceded all the work on the catalytic methods. Emde,³ by means of sodium amalgam, reduced cinnamyltrimethylammonium chloride to trimethylamine and propenylbenzene. He found this to be a reaction characteristic for quaternary ammonium compounds containing the cinnamyl radical. The corre-

$$[C_6H_6CH = CHCH_2N(CH_3)_3]X \xrightarrow{H_2}_{Na \cdot Hg}$$

$$C_6H_6CH$$
= $CHCH_3 + (CH_3)_3N + NaX$ 

sponding saturated compounds, [C₆H₅CH₂CH₂CH₂N(CH₃)₃]X and [C₆H₅CHClCHOHCH₂N(CH₃)₃]X, are stable under the same conditions. If the quaternary ammonium salt contains two cinnamyl groups, the products of the reaction are propenylbenzene and a cinnamyl-dialkylamine, which is stable until it is quaternized.

Benzyltrimethylammonium ehloride furnishes toluene and trimethylammonium ehloride and hydroxide are not

$$[C_6H_5CH_2N(CH_3)_3]Cl \rightarrow C_6H_5CH_3 + (CH_3)_3N + NaCl$$

affected by sodium amalgam. Dibenzyldimethylammonium chloride forms toluene and benzyldimethylamine. Tinnamylbenzyldimethylammonium chloride furnishes propenylbenzene and benzyldimethylamine, indicating that the cinnamyl-nitrogen bond is more easily cleaved under these conditions than is the benzyl-nitrogen bond.

$$\begin{bmatrix} \mathrm{CH_3} \\ \\ \mathrm{C_6H_5CH} = \mathrm{CHCH_2} - \mathrm{N} - \mathrm{CH_2C_6H_5} \\ \\ \mathrm{CH_3} \end{bmatrix} \mathrm{Cl} \, \rightarrow \,$$

 $C_6H_5CH$ = $CHCH_3 + C_6H_5CH_2N(CH_3)_2 + NaCl$ 

Hydrogenolytic cleavage of quaternary ammonium compounds has been used in the synthesis of methylpropylallylamine by the following sequence of reactions.6

$$(C_{6}H_{5}CH_{2})_{3}N \xrightarrow{CH_{3}I} \underbrace{(C_{6}H_{5}CH_{2})_{3}NCH_{3}II \xrightarrow{H_{2}} \underbrace{N_{a} \cdot H_{g}}}_{N_{a} \cdot H_{g}})$$

$$N_{2}I + C_{6}H_{5}CH_{3} + (C_{6}H_{5}CH_{2})_{2}NCH_{3} \xrightarrow{CH_{2}=CHCH_{2}I}$$

$$CH_{3}$$

$$CH_{3}CH_{2}CH_{2}CH_{2}$$

$$CH_{3}$$

$$CH_{3}CH_{2}CH_{2}CH_{2}I$$

$$CH_{3}CH_{2}CH_{2}CH_{2}I$$

$$CH_{2}CH_{2}CH_{2}CH_{2}I$$

$$CH_{3}CH_{2}CH_{2}CH_{2}I$$

$$CH_{3}CH_{2}CH_{2}I$$

$$CH_{3}CH_{2}CH_{2}I$$

$$CH_{3}CH_{2}CH_{2}I$$

$$CH_{2}CH_{2}CH_{2}I$$

$$CH_{3}CH_{2}CH_{2}I$$

$$CH_{2}CH_{2}CH_{2}I$$

An analogous reaction takes place in the reductive degradation of allocryptopine methosulfate (XI, R = CH₃) to methyltetrahydrocryptopine (XII, R = CH₃),⁸ and in the conversion of hunnemanine O-ethyl ether methosulfate (XI,  $R = C_2H_5$ ) to tetrahydromethylhunnemanine O-ethyl ether (XII,  $R = C_2H_5$ ).

⁶² Manske, Marion, and Ledingham, J. Am. Chem. Soc., 64, 1659 (1942).

auo-Cryptopine,  $R = CH_3$ Hunnemanine O-ethyl ether,  $R = C_2H_5$ 

Simultaneous Cleavage of Benzyl-Oxygen and Benzyl-Nitrogen Bonds (Table XIV). The simultaneous removal of benzyl groups attached to oxygen and to nitrogen offers nothing new in principle. Examples of these reactions are shown in Table XIV.

#### Cleavage of Benzyl-Sulfur Bonds

Debenzylation of benzyl thio ethers presents special problems. The sulfhydryl group in the product is likely to poison the ordinary catalysts and, hence, the usual catalytic procedures are not applicable. So-called "sulfactive" catalysts are employed in hydrogenolytic reactions, 63, 64 but their use is not restricted to the removal of benzyl groups. Raney nickel as usually prepared contains appreciable amounts of hydrogen and will not only split thio ethers but will remove a sulfur atom, and such desulfurization is not limited to benzyl thio ethers. 65, 66 Catalytic procedures limited to the hydrogenolysis of benzyl-sulfur linkages have not been described.

Chemical methods, however, are available for S-debenzylation. They are extensions of the chemical methods used for removing the carbobenzyloxy group described on p. 275. Sodium in liquid ammonia reacts with carbobenzyloxycysteine to remove the carbobenzyloxy group and does not affect the sulfhydryl group. In these experiments the cysteine was not isolated but was oxidized to cystine, which was isolated in almost quantitative yield. When S-benzylcysteine was treated with sodium in liquid ammonia, debenzylation took place; the debenzylated product was oxidized, and cystine was isolated in a yield of 80%. The benzyl group appears not as toluene but as bibenzyl. Similar procedures

⁶¹ Signaigo, U. S. pat. 2,402,686 [C. A., 40, 5766 (1946)].

⁴ Farlow, Hunt, Langkammerer, Lazier, Peppel, and Signaigo, J. Am. Chem. Soc., 70, 1392 (1948).

Bougault, Cattelain, and Chabrier, Compt. rend., 208, 657 (1939).

Mozingo. Wolf, Harris, and Folkers, J. Am. Chem. Soc., 65, 1013 (1943).

 $C_6H_6CH_2SCH_2CH(NH_2)CO_2H \xrightarrow{Na}$  Liquid NH₃

 $C_6H_6CH_2CH_2C_6H_6 + HSCH_2CH(NH_2)CO_2H$ 

 $\int O_2$ 

SCH2CH(NH2)CO2H SCH₂CH(NH₂)CO₂H

have been used for the preparation of homocystine,67 dideuteromethionine and tetradeuterocystine,  68  and  $\alpha$ -amino- $\beta$ -mercaptobutyric acid.  69 

# EXPERIMENTAL CONDITIONS AND CATALYSTS

Various palladium catalysts are described by Mozingo; 70 palladium black is prepared according to the directions of Tausz and Putnocky; 71 platinum black is described by Feulgen; 72 platinic oxide by Adams, Voorhees, and Shriner; 73 Raney nickel by Covert and Adkins. 74 Workers experienced with catalytic procedures need not be reminded that there are many modifications in the methods of preparing catalysts, especially those derived from the noble metals, and that there are still some imponderables in the process.

Catalytic reductions are usually carried out in the standard apparatus,75 and in the absence of side reactions the course of hydrogenolysis parallels the drop in pressure of hydrogen. The choice of solvents is large. The effects of higher pressures have not been assayed, but generally it may be said that with palladium and platinum no high pressures are required and room temperature is usually adequate.

# EXPERIMENTAL PROCEDURES

o-Tolylglucoside from Salicin.  18  In a microhydrogenation apparatus  76 is placed 0.25 g. of salicin in 25 ml. of water containing a trace of hydrochloric acid; 0.05 g. each of platinum black and palladium black are added. Absorption of hydrogen stops after one mole is taken up, in

F Patterson and du Vigneaud, J. Biol. Chem., 111, 393 (1935).

⁶³ Patterson and du Vigneaud, J. Biol. Chem., 123, 327 (1938).

⁶⁹ Carter, Stevens, and Ney, J. Biol. Chem., 139, 247 (1941).

⁷⁰ Mozingo, Org. Syntheses, 26, 77 (1946).

⁷¹ Tausz and Putnocky, Ber., 52, 1576 (1919).

Adams, Voorhees, and Shriner, Org. Syntheses Coll. Vol., I, 463 (1941).

⁷⁴ Covert and Adkins, J. Am. Chem. Soc., 54, 4116 (1932).

Adams and Voorhees, Org. Syntheses Coll. Vol., I, 61 (1941). ⁷⁶ Hyde and Scherp, J. Am. Chem. Soc., 52, 3359 (1930).

are then combined with the organic layer and dried; the solvent is removed at reduced pressure and the residue fractionated in vacuum; the p-dimethylaminotoluene distils at 77-79°/6.5 mm. and weighs 10.5 g. (78%).

p-Aminodiphenylmethane from p-Aminobenzophenone. In a 2-l. Soxhlet flask is placed 136 ml. of 1.2 M lithium aluminum hydride in diethyl ether (7 equivalents per mole of ketone), diluted with 200 ml. each of benzene and dibutyl ether. The contents of the flask are heated to boiling (80°), and then a Soxhlet extraction apparatus is mounted on the flask, the thimble of the apparatus being charged with 13.8 g. (0.07 mole) of p-aminobenzophenone. Vigorous refluxing at 80° is maintained for one hour. The reaction mixture is cooled and carefully hydrolyzed with 200 ml. of 5% sodium hydroxide solution. The organic phase is separated, and the aqueous suspension is extracted with five 200-ml. portions of diethyl ether. The combined extracts, with the organic layer, are freed from solvents at reduced pressure. The residual viscous red oil is extracted repeatedly with hexane to yield 7.3 g. (57%) of a yellow oil which crystallizes on cooling with acetone and solid carbon dioxide. After drying, the p-aminodiphenylmethane melts at 34-35°.

The residue from the hexane extractions is a dark gum which, after crystallization from benzene, yields 2.1 g. (15%) of crude p-aminobenzhydrol, m.p. 108-112°; on repeated crystallization from water the product melts at 116-117°.

Dihydromorphine from Benzylmorphine." Twenty-five grams of benzylmorphine hydrochloride is suspended in water and shaken in a hydrogen atmosphere with palladium-charcoal catalyst. Two moles of hydrogen is taken up. The catalyst is filtered, and from the filtrate are isolated toluene and dihydromorphine, the latter being recovered in quantitative yield by volatilizing the toluene and the water.

Toluene and Butanol from n-Butyl Benzyl Ether. In a copper liner inside a steel bomb is placed 46 g. of n-butyl benzyl ether and 2.5 g. of Raney nickel. Hydrogenation is carried out at 175° and 150-200 atm. After one and one-half hours 93% of the ether has been converted to toluene and butanol.

5-Phenyl-1-pentanol from 2-Phenyltetrahydropyran. Nine grams (0.056 mole) of 2-phenyltetrahydropyran is dissolved in 40 ml. of acetic acid solution containing 2.5% of 60% perchloric acid; 100 mg. of palladium-charcoal catalyst (5%) is added, and the mixture is reduced in the ordinary apparatus at 3 atm. Reduction is complete in thirty-five minutes. The catalyst is removed, the filtrate is poured into 10% sodium hydroxide solution, and 5-phenyl-1-pentanol is extracted with

ether or with tetrachloroethane and distilled, b.p. 142-148°/10 mm.; yield 72%.

(+,-)-Phenylalanylglycine from β-Phenyl-α-benzyloximinopropionylglycine. Four grams (0.0123 mole) of β-phenyl-α-benzyloximinopropionylglycine is dissolved in a solution of 150 ml. of water and 2.5 ml. of concentrated ammonium hydroxide. The hydrogenation is carried out at 3 atm., using 3.5 g. of palladium catalyst (10%), and requires about two hours. The catalyst is then removed, and the filtrate is evaporated to dryness at reduced pressure and over a steam bath. The residue is triturated with methanol and washed with ether. The product, which is the dihydrate, weighs 2.4 g. (87%). It may be completely dried over phosphorus pentoxide to furnish (+,-)-phenylalanylglycine, m.p. 273–275° (dec).

3-Glyceraldehyde Phosphate from Benzylcycloacetalglyceraldehyde Phosphate. In an apparatus which assures an atmosphere of pure hydrogen is placed 0.6 g. of palladium catalyst and 10 ml. of acetic acid which has been distilled from chromic acid. In a special bulb is placed 1.3 g. of pure benzylcycloacetalglyceraldehyde phosphate. The apparatus is shaken to saturate the catalyst; then the special bulb is inverted to add the substrate to the reaction mixture and shaking is resumed. Hydrogenolysis is complete in thirty to forty minutes at room temperature. The hydrogen in the apparatus is replaced by air, the mixture is removed and filtered, and the filtrate is concentrated at 30° at reduced pressure. The residue is washed on the centrifuge with one 4-ml. and with two 2-ml. portions of water; the undissolved substance is unchanged starting material. The combined aqueous washings are again concentrated at 30° to a syrup; final desiccation is achieved at 0.05 mm. The product is purified by washing on the centrifuge with methanol.

Barium D-Glucose-6-phosphate from 1,2-Isopropylidene-D-glucose. Dibenzyl chlorophosphonate, from 13.1 g. of dibenzyl phosphite, in 50 ml. of dry chloroform is added dropwise over a period of seventy-five minutes to a stirred solution of 11.0 g. of 1,2-isopropylidene-D-glucose in 100 ml. of pyridine at -10°. The mixture is allowed to warm to room temperature as stirring is continued and is then allowed to stand overnight. It is evaporated at reduced pressure, and the residual syrup is taken up in chloroform, washed with dilute sulfuric acid, then with water, and dried over anhydrous sodium sulfate; the solvent is evaporated. The residue is dissolved in ethanol, and the solution is heated to reflux for thirty minutes with 5 g. of Raney nickel to remove possible catalyst poisons. The solution is filtered and hydrogenated with a mixed catalyst, 0.5 g. of palladium oxide and 1.0 g. of palladium-charcoal

⁷⁹ Fischer and Baer, Ber., 65, 337 (1932).

(10%), until no more hydrogen is taken up. The solution is filtered to remove catalyst. The isopropylidene group is removed by acid hydrolysis. p-Glucose-6-phosphate is isolated as the barium salt,  $[\alpha]_D^{30}$ + 11.8°. The yield is 9 g. (42%).

2-Glycerol-β-D-glucoside from 1,3-Benzylideneglycerol-β-D-gluco-Benzylideneglycerol-β-p-glucoside, 1.25 g., is dissolved in 100 ml. of absolute ethanol and shaken with 0.9 g. of palladium black in an atmosphere of hydrogen. After an hour the hydrogen uptake ceases and glycerol- $\beta$ -p-glucoside precipitates. It is filtered with the catalyst, from which it may be removed by dissolving in water. Evaporation of the aqueous solution leaves 0.9 g. (97%) of crystalline 2-glycerol- $\beta$ -Dglucoside, m.p. 165°.

Phenylacetic Acid from Acetylmandelic Acid.43 Two grams of acetylmandelic acid is dissolved in 10 g. of tetralin, and several grams of palladium-barium sulfate is suspended in the solution. The suspension is heated to 215°, the refluxing temperature of the solvent, and hydrogen is passed through for six hours, entering at the bottom of the boiling mixture. The mixture is then cooled and the catalyst removed. The phenylacetic acid is extracted with sodium carbonate solution, from which it is recovered by acidifying with hydrochloric or sulfuric acid. Crystallization from water yields the pure acid, m.p. 76° (60%).

L-Glutamylglycine Ethyl Ester from Carbobenzyloxy-L-glutamylglycine Ethyl Ester. 81 A solution of 8.2 g. of carbobenzyloxy-I-glutamylglycine ethyl ester in about 50 ml. of ethanol containing 2 ml. of glacial acetic acid is shaken with platinum black catalyst. After hydrogen absorption has ceased, the catalyst is removed and the solution evaporated; the residue is evaporated repeatedly with ethanol. The spongy mass which precipitates from ethanol on the addition of ether weighs 4.1 g. (80%). L-Glutamylglycine ethyl ester melts at 151°.

Diglycyl-L-cystine from Dicarbobenzyloxyglycyl-L-cystine.82 To a stirred solution of 25 g. of dicarbobenzyloxyglycyl-L-cystine in 250 ml. of liquid ammonia are added small pieces of sodium until a blue color appears. The ammonia is then allowed to volatilize spontaneously, and the residual traces of ammonia are removed by evacuating the container for several hours on the water pump. The residue is taken up in cold water, and dilute sulfuric acid is added until the solution is acid to litmus. The glycylcysteine is precipitated with mercuric sulfate reagent, washed several times with water, and centrifuged. The complex is decomposed with hydrogen sulfide, and the precipitation with mercuric sulfate is

⁸¹ Bergmann, Zervas, and Fruton, J. Biol. Chem., 111, 225 (1935).

⁸² Greenstein, J. Biol. Chem., 128, 241 (1939).

repeated. The final solution is made slightly alkaline with barium hydroxide solution, and the precipitated barium sulfate is removed by centrifuging. A few crystals of ferric oxide are added to the solution, and air is bubbled through it until the test with sodium nitroprusside shows the sulfhydryl group to be absent. The solution is heated with decolorizing charcoal, and the barium is precipitated quantitatively by the addition of sulfuric acid. The filtered solution is evaporated almost to dryness at reduced pressure. On addition of ethanol to the concentrate, the oxidized peptide, diglycylcystine, precipitates in gelatinous form. The mass is taken up in water and precipitated with ethanol, the process being repeated several times. After the last precipitation the mass is heated. It dissolves in the adhering ethanol and the peptide crystallizes from the hot solution in long prisms. The yield is 8.0 g. (57%), m.p. 232° (dec.),  $[\alpha]_D^{24} - 108$ ° for 75% solution in 0.1 N hydrochloric acid.

Di-n-hexylamine from Benzyldi-n-hexylamine. A solution of 27.0 g. of benzyldi-n-hexylamine in 30 ml. of glacial acetic acid is shaken with 0.4 g. of platinic oxide in an atmosphere of hydrogen at 70°. After six hours the reduction is complete. The catalyst is removed, the filtrate is made strongly alkaline, and the di-n-hexylamine is extracted with diethyl ether. The extract is dried and fractionated; the amine distils at 110°/14 mm. The yield is practically quantitative.

Dialkylamines from Benzyldialkylamines.⁵² The benzyldialkylamine, as free base or salt, is dissolved in twice its weight of glacial acetic acid, and platinum oxide catalyst, usually 1% of the weight of the amine, is added. Hydrogenation is carried out at 65–75° and 3 atm. Eight hours or less are required for reduction. The reaction mixture is diluted with methanol, the catalyst is removed by filtration, and excess hydrochloric acid is added to the filtrate which is concentrated at reduced pressure. To liberate any acetylated amine, the residue is digested on the steam bath with concentrated hydrochloric acid, 50 ml. for 0.1 mole amine, for several hours. Evaporation of the liquid leaves the amine hydrochloride, which may be purified by crystallization from an appropriate solvent; or the residue may be treated with alkali to liberate the free secondary amine, which may then be distilled.

2,3,5-Trimethylphenol from 2-Dimethylaminomethyl-3,5-dimethylphenol.⁶¹ A solution of 18 g. of 2-dimethylaminomethyl-3,5-dimethylphenol in 200 ml. of dioxane is hydrogenated in the presence of 7.5 g. of copper chromium oxide for four hours at 165° and 177 atm. The catalyst is removed and the dioxane distilled. The residue, after acidification with a small amount of hydrochloric acid, is distilled with steam to

furnish 8 g. (58%) of 2,3,5-trimethylphenol. The product, crystallized from petroleum ether, melts at 93°.

1-(3,4-Dihydroxyphenyl)-2-amino-1-butanol from  $\alpha$ -Benzhydrylamino-3,4-dibenzyloxybutyrophenone.14 To a solution of 28.9 g. (0.1 mole) of  $\alpha$ -benzhydrylamino-3,4-dibenzyloxybutyrophenone hydrochloride in 150 ml. of absolute methanol, 0.5 g. of palladium sponge is added. The mixture is shaken with hydrogen at 55-70° and 3 atm. until 3 moles of hydrogen is taken up. The catalyst is removed, the toluene and the diphenylmethane are extracted with ether, and the aqueous layer is decolorized with charcoal and further hydrogenated with fresh catalyst until a fourth mole of hydrogen is taken up. The catalyst is again removed and the filtrate taken to dryness under reduced pressure. The residue is dissolved in absolute ethanol and again decolorized; then acetone and dry ether are added until precipitation is complete. The product weighs 14 g. (60%) and melts at 199–200° (dec.).

Benzylhydrazine from 1,1-Dibenzylhydrazine. A solution of 4.1 g. of 1,1-dibenzylhydrazine in 50 ml. of absolute ethanol is hydrogenated with 400 mg. of palladium oxide. After hydrogen absorption ceases, the catalyst is removed and dry hydrogen chloride is led into the filtrate, whereupon 2.7 g. (88%) of benzylhydrazine hydrochloride precipitates. The product may be crystallized from ethanol.

Benzyldimethylammonium Chloride from Dibenzyldimethylammonium chloride.3 Fifteen grams of dibenzyldimethylammonium chloride is dissolved in 50 ml. of water. Over a period of two days 50 g. of 5%sodium amalgam is added in small portions at room temperature. There is little evolution of gas, the solution becomes turbid, and after several hours an appreciable oily layer accumulates on the surface. On the second day the aqueous solution becomes clear, and the addition of more sodium now causes a vigorous evolution of gas. The liquid is decanted from mercury and extracted with ether; the aqueous layer contains a very small amount (about 0.1 g.) of the unchanged quaternary ammonium salt. From the ethereal extract the amine is removed with dilute hydrochloric acid. Concentration of the acidic extract leaves 9.0 g. of benzyldimethylammonium chloride (91%). Toluene may be recovered from the ether layer.

D-Homocystine from S-Benzyl-D-homocysteine.83 A solution of 6.4 g. of S-benzyl-D-homocysteine in 40 ml. of liquid ammonia is treated with a slight excess of metallic sodium. The ammonia is allowed to evaporate spontaneously, and the residue is dissolved in 60 ml. of water. One-tenth gram of hydrated ferric chloride is added, and air is passed through the solution until the test with sodium nitroprusside for free sulfhydryl

⁸³ du Vigneaud and Patterson, J. Biol. Chem., 109, 97 (1935).

groups is negative. The precipitated ferric hydroxide is removed by filtration, and the clear filtrate is made neutral to litmus with dilute hydrochloric acid. Pure p-homocystine precipitates; 2.85 g. (75%); after recrystallization from water the product melts at 281–284° (dec.).

 $\alpha$ -Amino- $\beta$ -mercapto-n-butyric Acid from  $\alpha$ -Amino- $\beta$ -benzylmercapto-n-butyric Acid.⁶⁹ Fifteen grams of  $\alpha$ -amino- $\beta$ -benzylmercapton-butyric acid is dissolved in 250 ml. of liquid ammonia and treated with small pieces of metallic sodium slightly more than two equivalents being necessary to produce a permanent blue color. Enough ammonium chloride is then added to discharge the color, plus 7 g. additional. The ammonia is allowed to evaporate, the final traces being removed at reduced pressure. To the residue are added 250 ml. of ether and 5 ml. of concentrated hydrochloric acid; the mixture is stirred and heated on the steam cone for several minutes. The ether is decanted, and the residue is again extracted with ether. The subsequent operations are carried out in an atmosphere of nitrogen. The residue is extracted with three 100-ml. portions of warm absolute ethanol containing a few drops of concentrated hydrochloric acid, and the combined extracts are taken to dryness under reduced pressure. The residue is dissolved in 80 ml. of absolute ethanol, and 800 ml. of anhydrous ether is added. The solution is cooled overnight, and the precipitate removed, washed with ether, and dried, yielding 9.8 g. of  $\alpha$ -amino- $\beta$ -mercapto-n-butyric acid hydrochloride. This is dissolved in 300 ml. of ethanol, and 3.8 ml. of concentrated ammonium hydroxide is added; on cooling, 6.4 g. (71%) of pure amino acid is obtained, m.p. 203-204° (dec.).

#### TABULAR SURVEY

In the seventeen tables that follow are listed examples of the reductive cleavage of benzyl groups. As indicated earlier, it is not possible to guarantee the completeness of the tables because many examples of the reaction are subordinated to other aspects of the articles in which they appeared. The survey of the literature was carried to July 1950.

TABLE I

BENZYL ALCOHOLS

Dofor	•	Rapid 16 2.5 hr. 84	2 hr. 85		6 d. 32			20 min. 18		
Pres-	sure atm. Ti 3 Ra	3 Ro 220-240 2.	375 2		9	1		1 20		
		25 185	086	007	8	١		25		
	Solvent	Ethanol Ethanol Abs. CH ₃ OH	į	Dioxane	$(C_2H_6)_2O$		i	Н20		
Benzyl Alcohols	Catalyst	Pd-charcoal Pd-charcoal Copper chromium	oxide	Copper chromium Dioxane	oxide LiAlHa	-	Colloidal Pt	Pt or Pd black		
BENZYL	Yield %	Quant. Quant.	3	84	ri Cr	3	1	Quant.		
	Product Isolated		p-CH ₃ OC ₆ H ₄ CH ₃	3,4-(CH2O2)C6H3CH3		0-H2NC6H4CH3	C6H11O5-0-C6H4CH3	(o-tolylglucoside)	(o-tolylglucoside)	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	South Co.	4		3 ALCHONC, HOCHOOH		o-H2NC6H4CH2OH	C.H.,O. — O.—C.H.CH2OH C.H1105.—O.—C.H4CH3	(salicin) (o-tolylglucoside)	(salicin)	111111111111111111111111111111111111111

Note: References 84-165 are listed on pp. 325-326.

#### ORGANIC REACTIONS

Refer-ence 16

Pres-sure atm.

Tem-pera-turo °C.

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ALC
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5

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cuce	16	16	27 27	23	27	1 12 8	16		98	87		 8 8		
Time	Moder-	Moder-	10 min. 0.5 hr.	i	١	1 1	Moder-	3	2.5 hr.	2 br.	3.5 br.			
eure atm.		8			-				147	170	3.5	2.5		
turo B		25	80-90	80-00	80-90	80-90	25 25		160	200	25	25. 45.		
	lycat	Ethano.				CH3CO2H + HC104 CH3CO2H + HC104	CH ₃ CO ₂ H + HO ₁ O ₃ Ethanol + HO ₁	Ethanor	ď,	, n	а Н2О	CH3CO2H + H2SO4 CH3CO2H + HCIO4	CH1CO1H + H2SO1	
	Catalyst	Quant. Pd-charcoal	Pd-charcoal		Post-Pd	Pd-BaSO4		Pd-charcoal	•	Copper chromium 1120	Copper ebromium	PĂ I		
	Yicld %	Juant.	i	08-09 08-09	08-09	8	08-09 80-80 80-80	18		90	90	•	88 75	
) \$	<del>-</del>	ict Isolated	Cancing + Cancing		CellCil(Cilt)NII2 CellCell(Cilt(Cilt)NII2	MOHO HOUSE	O,II,CH1CH1CH(C2H2)NH2 P-11OC,H1CH1CH(C2H3)NH2	Conscionants Conscionants	3,4-(01130)-4,6	CHCHACH2CH2CO2Na	p-Closs CH(CH3)CH2CH2CO2Na	PCH,O,H,G(OH)(CH3)CH,CH2CO,Na p-CH3Co,HQC,HQC,HQC	Conschoolsh Conschoolsh	C ₁ II ₁ CII ₂ CO ₂ C ₂ II ₁
		Substance Reduced	CinchonGins	Callactionicity off	Cellicitoticit(Cilis)NIIs	p-CII,C411,CIIOIICII(C2II6)NII?	C. II, CHOHCH(C. II,) NHCH3	P-011,0C6,11,C11011C11(C2,115,N1112	3,4.(CII3O),C4II3CIIOIICN		$\beta \cdot C_{10}I1_7C(OI1)(CII_3)CII_2CII_2CO_2N\alpha$	p-CH1,C4H,C(OH)(CH13)CH12CH2CO:1	CelliCITOIICO1II	Ctricitoricostri Ctricitoricoscitis

	HY	DRO	)GE	NOLYSIS OF	BE
88 88 88 88 89	83 83	8 8 8	So So	ដ ដ ដ ដ ដ	
8 hr. 1.5 hr. — 1.5 hr. 6 min.	17 min.	1.1	l	2.5 hr. 2.5 hr. 2 hr. 1.5 hr. 1.5 hr.	
2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	61 63	61 61	1	ना ना च च च	
25 100 25 60 25	25	22 22		25 25 25 25 25	
CH ₃ CO ₂ H + HClO ₄ CH ₃ CO ₂ H CH ₃ CO ₂ H + HClO ₄ CH ₃ CO ₂ H + HClO ₄ CH ₃ OH + H ₂ SO ₄	CH3OH + HCI	CH3CO2H + H2SO4	Gracos Benzene	Abs. ethanol Ahs. ethanol Abs. ethanol Abs. ethanol Abs. ethanol	
Pd black Pd black Pd black Pd black Pd-charecoal	Pd-charcoal	Pd-charcoal Pd-charcoal	Pd-charcoal Pd black	Pd-charcoal Pd-charcoal L. Pd-charcoal Pd-charcoal Pd-charcoal	
90 88 77 92	S	88 57	5 8 8	60 60 Quant. 80 75	
C6H1CH2CO2C2H1 C6H1CH2CO2C2H5 p-C2H1CH4CH2CO2C2H5 p-C2H1C6H2CH2CO2C2H5	C ₆ H ₆ CH ₂ CH ₂ NH ₂	Chichichinhi Chichich	Conf. CH2CH2NH2 Conf. CH2CH2NH2	CcHcCN p-CH ₃ OC ₆ H ₄ (CH ₃ ) ₃ CH ₃ p-CH ₃ OC ₆ H ₄ CH ₂ CH(CH ₃ ) ₂ C ₆ H ₅ CH ₂ CH ₃ C ₆ H ₅ C ₆ H ₅ CH ₂ CG ₆ H ₅ C ₆ H ₅ CH ₂ CG ₆ H ₅	l on op. 325-326.
Chuschohcosch Chichtohcosch Chischtohcosch	P-C2H (CH (OCOC2H)) CN	Of H COCOCAH DON	Chhich (CCOCH3) CN Chich (CCOCH3) CN	Chronocation Chronocation p-Chronocation Chronocation Chronocation Chronocation Chronocation Chronocation Chronocation	225-326.

Note: References 84-165 are listed on pp. 325-326.

HYDROGENOLYSIS OF BENZYL GROUPS
22 20 20 20 20 30 90 90 90 90 90 16 16 16 16 16 17 27 27 27 27 27 27 27 27 27 27 27 27 27
45-90 min. 45-90 min. 45-90 min. 45-90 min. 45-90 min.  Moderate
**************************************
Ethanol CH3OH CH3OO CH3OO2H + HCIO4 CH3CO2H + HCIO4
d-charcoal
- Pd- Quant. Pd- Quant. Pd- Quant. Pd- 75 Pd 70 Pd 86 Pd- 86 Pd 87 Pd 71
m-CH ₃ OC ₆ H ₄ CH ₂ C ₂ H ₆ m-HOC ₆ H ₄ CH ₂ C ₂ H ₆ m-HOC ₆ H ₄ CH ₂ C ₂ H ₆ 3.4-(HO) ₂ C ₆ H ₃ CH ₂
m-CH, OC&H, COC2H6  m-HOC, MICOC2H6  m-HOC, MICOC2H6  3,4-(HO), 2-641; OCC2H6  3,4-(HO), 2-641; OCCH, CH3, 13,4-(H3), 2-641; OCCH, CH3, 13,4-(H3), 2-641; OCCH, CH3, 13,4-(H3), 2-641; OCCH, CH3, 13,4-(H3), 2-641; OCCH, CH3, CH3, CH3, CH3, CH3, CH3, CH3,

Note: References 84-165 are listed on pp. 325-326.

Reduction to the cathinol was rapid; reduction of the earbinol was slow and incomplete. · Reduction to the carbinol was rapid; reduction of the carbinol was slow. The reduction was rapid until half completed, then slow.

The state of the s

# TABLE IV-Continued

		ORGAN	IC REACTION	NS			
	Refer- ence 27	;	ă.	93	16	អ	ព ព
	Tine 1	:	5-10 br.	5-10 hr.	<u>i</u>	1 14.	1 hr.
	Pres-		ı	130	1	-	4 4
Теш-			8	0110	81	ដ	<u> </u>
	Solvent CH1CO2H + HClO4		NaOH + H ₂ O	H ₂ O	Abs. ethanol	Abs. ethanol	Abs. ethanol Abs. ethanol
	Catalyst Pd-BaSO4		Copper chromium oxide	Copper chromium	Copper chromium orido	Quant. Pd-charcoal Aba ethanol	Quant, Fd-charcoal Abs. ethanol Quant, Fd-charcoal Abs. ethanol
	Yield %		99	18	<b>ତ</b>	Quant.	Quant. Quant
Ketones	Product Isolated H H	C NH2	CH ₂ CH ₂ CH ₂ CH ₂ CO ₂ Na	a-CloH;CH(CH;)CH(CH;)CO;N3	Z HZ	C ₆ H ₅ (CH ₂ ) ₁ C ₆ H ₅	P-CH ₂ C ₆ H ₄ (CH ₂ ) ₃ C ₆ H ₅ C ₆ H ₅ (CH ₂ ) ₃ C ₆ H ₄ OCH ₂ -p
	Substance Reduced	HON-SO	COCH2CH2CO2NA	/ α-C ₁₀ H ₇ COCH(CH ₃ )CH(CH ₂ )CO ₂ N ₃		C,H,COCH=CHC,H,	P-CH ₂ C ₂ H ₄ COCH=CHC ₂ H ₅ C ₂ H ₅ COCH=CHC ₅ H ₄ OCH _{5-P}

HYD)	ROGENOLYSIS OF	BENZYL GROUPS
233 99 99 99 99	33 33 11	Reference 19 19 19 19 19 19 19
1 hr. 1.5 hr. 3.5 hr. 3 hr. 25 min. 40 min. 1.25 hr. 8 hr.	1 hr. 1 hr. 11 d.	Time 30 min. 1.5 hr. 30 min. 1.5 hr. 30 min. 7 hr.
3.5 3.5 3.5 2.6 40 11.1 2.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8 12.6 8	2.5	Pres- sure atm. 150-250 150-260 150-250 150-250 150-250
55 50 50 50 50 50 50 50 50 50 50 50 50 5	8 8 8 8 8	Temper- ature °C. 160  175 25 25 26 160 100
. Н ₂ SO4		Solvent
Abs. ethanol Abs. cthanol CH ₃ CO ₂ H + H ₂ SO ₄ CH ₃ CO ₂ H + H ₂ SO ₄ CH ₃ CO ₂ H	СН 3ОН (С3H5)2О (С2H5)2О (С2H5)2О	Catalyst Raney Ni H2PtCle Raney Ni Raney Ni Raney Ni Pd-charcoal Raney Ni Raney Ni
Pd-charcoal Pd-charcoal Pd black Pd black Pd-charcoal Pd-charcoal Pd-charcoal Pd-charcoal	Pd Black-S LiahH4 LiahH4 LiahH4	Yield % C C 64 Ran H2 92 Ra 80 Ra 60 Ra 72 Ra 72 Ra 71
94 44 44 44 44 44 44 44 44 44 44 44 44 4	57 32 46	ಭ
	ę.	TABLE V BENZYL ETHERS Roduct Isolated 50H I
CeHs(CH2)3CeH3(O2CH2)-3,4 CeHs(CH2)1CeH1s p-CH3(OCH4,CH2,CH2,NH2 3,4-(CH3,O)2CH3,CH3,CH3,NH2 p-CH3(CH2)1CO2H p-CH3(CH2)1CO2H p-CH3(CH2)1CO2H c-Ch4(CH2)1CO2H c-Ch4(CH2)1CO2H	CH4CoH6 CH4CoH6 P-H2NCH4CH2CH8 p-H2NCH4CH3CoH4CH5P p-CH30CcH4CH2CoH4CH5P p-CH30CcH4CH2CoH4CH5P	B1
C ₆ H ₅ COCH=CHC ₆ H ₃ (O ₉ CH ₂ )-3,4 C ₆ H ₅ COCH=CHCH=CHC ₆ H ₆ p-CH ₃ OC ₆ H ₄ COCN 3.4-(CH ₅ O) ₂ C ₆ H ₃ COCN p-CH ₃ OC ₆ H ₄ COCH ₂ CO ₂ H p-CH ₃ OC ₆ H ₄ COCH ₂ CO ₂ H c ₆ C ₆ H ₆ COCH ₃ CO ₂ H C ₆ H ₆ COHCOCG ₆ H ₆ CO ₃ H	CH3COOCH3  ChBCOCH3  P-H2NCAH4COCh4NH2-p  p-R3NCAH4COCh4ANH2-p  p-CH3Coch4COCh4ACH3-p  Node: References 84-165 are lieted on pp. 325-326,	Substance Reduced CeHeOH2OCH3 CeHGCH2OCH3 CeHGCH2OCH03 CeHGCH2OCH(GH3)2CH3 CeHGCH2OCH(GH3)2CH5 CeHGCH2OCH(GH3)2CH6 CeHGCH2OCH(GH3)2CH6 CeHGCH2OCH294CH3

Note: References 84-165 are listed un pp. 325-326.

Refer-

Temper- Pres-Sure

## TABLE V-Continued

## BENZYL ETHERS

	And the state of t	Yield %	Catalyst	Solvent	ature °C.	sure atm.	Time	ence
Substance Reduced	Product Boures C.H.OCH2CH2OH	2 2 5	Raney Ni	li	175 150	150-250 150-250	4 hr. 1,3 hr.	19
$C_0H_0CH_2CH_2CU_2H_0$ $C_0H_0CH_2OCH_2CC_0H_0$ $2\cdot 3(CH_1O).C_0H_1CH=CH(CH_2)_0OCH_2C_0H_0$	C6H2OCH2OH2OH4CH2)7OH	97 Quant.	Pd black Pd-charcoal	CH1CO2H CH1CO2H	នន	2-3		38
C,U,CH,CH,CH,CH,CH,CH,CH,CH,CH,CH,CH,CH,CH,	$c_{eH_bOH} = c_{eH_{11}OH}$	Quant.	Raney Ni	l	18	150-250	24 min.	1
	HO,H2O0,H2-0	Quant.	Pd-charcoal	1 1	ន ន	1 150-250	ij	11
ø-CH ₂ OC¢H4OCH2C¢H8 CH-CaH4OCH2CAH8	o-CH ₁ C ₆ H ₁ OH	8 2	Raney Ni	! <b>!</b>	150	150-250	_	19
m.CH1CaH4OCH2CaHs	m-CH ₃ C ₆ H ₄ OH + m-CH ₃ C ₆ H ₁₀ UH	: 8	Raney Ni	l	150	150-250		£ 13
P-CH3C6H4OCH2C6H8	p-CH1CtH1OH + p-CH1CtH10Ct o-CH1O2CCcH1OH	11	Raney Ni	l	120	150-250	24 min.	2
9-CH302CC6H40CH2C6H3		1	2	ł	22	-	l	18
C.H.CH.OC.H.10.	CeH12Os (glucose)	Ì	1 6	l	ĸ		l	138
C.H.CH.OCAH110s	C6H12O6 (glucose) + C6H11O6OCH2C6H11	1 %	P. black	Dil. HCl	23	1	90 min.	28
C,H,CH2OC,H11Os	C ₆ H ₁₂ O ₆ (glucose) C ₆ H ₁₂ O ₆ (glucose)	66	Pd black	H20	ន	-	3 hr.	<b>5</b> 2
Сенесначения		8	Pd sharmal Ethanol	Ethonol	13	-	l	34
# COOK	H,C	3		+ toluene				
Ms CERCE	но							
CH ₃	$CH_{\eta}$						,	7,
	HO	ł	ı	1	1	l	1	3
C, H, CH2O	HO NO							
C, H ₅ CH ₂ O	<u> </u>							
ĊH2	CH ₂							
OCH2	-CCH2					,		:
C.H.CH.CH.OC17H18O2N (benzylmorphine)	C17H21O3N (dihydromorphine)	Quant.	Quant, Pd-charcoal H20	п₂0	£	<b>-</b> l	ll	96
P-C,HCH2OC,H,COCH2N(CH3)COC,HS	p-HOC6H,COCH2N(CH3)COC6Hs	1	D.	l				

97

	]	HYDROC	ENOL	ASIS OF BEV	NELL	GROOTS		
97	25	34	43	66	17	001 008	16	12
1	1	10 min. 3 min.	1	9 min.	1 1	1 1		30 min. 20 hr.
1	6. 10.	- <del>-</del>	-	-	1 1	1 -	က	150-250
i	22	82 83	ដ	22	<b>i</b> 1	<del>গ্</del> ল	22	125 40–50
1	1	Abs. CH ₁ OH Abs. CH ₁ OH-	70% CH1CO2H	сн,сол	Ethanol	CH3CO2H † Ethanol	l Ethanol	60 Raney Ni 60 Pt black CH ₃ CO ₂ H
Pt02	Pt02	Pd-charcoal Pd-charcoal	Pd	Pd	Colloidal Pt Na +	etnamor Pt	Pd-charcoal Ethanol	Raney Ni Pt black
1	1	96 ) Quant.	1	ક્ર	Good	Quant.	Quant.	09
Dibydronsphthalene derivative; no debenzylation	Dibydronsphthalene derivative; no debenzylation	3-CH ₂ O-4-HOC ₆ H ₃ CHOHCH(NH ₂ )CH ₃ 3.4-(HO) ₂ -C ₃ H ₃ CHOHCH(NH ₂ )CH ₃	сно	2CHOH CH2OPO3H2 CH2OCH3 CCHOH CCHOH	CcH 1206+C9H18O2 (tetrahydrodesoxyaucuhigenin) Discetoneglucose	Monoscetoneglucose  CH ₂ CO ₂ CH ₂ CH[CH(OCOCH ₃ )] ₄ CHOCH  CH ₂ CO ₃ CH ₂ CH[CH(OCOCH ₃ )] ₄ CHOCH	P-CH1OC6H4CH1	Hydrogenated \$\textit{\textit{P-naphthols}}\$\$CH_2OH(CHOCOCH_1\textit{ACH_2OH}\$\$
CHOHCH ₂ N(C ₃ H ₇ -n) ₂	CHOHOLH Pn)2	OCH-CoH. CH-COCH-COCH-OCH-CH-CH-COCH-CH-COCH-COC	3,4-(c,H,CH,CH,CHORDICA3 BaOgPOH2C—CH—CH—OCH2C,Hs	C.H.GH.2O—CH—CH—CH-OPO.Ba. CH.OH.5C—CH—CH—OCH.2C.H.6 O O O C.H.GH.2O—CH—CH—CH-OCH.1	CgH11OsOCgH13O3 (aucubin) Diacetonebenzylglucose	Diacetonebenzylglucose  CH2OCH2CH6  CH3CCACH2CH[CH(OCOCH3)]sCHOCH	 CH1OCH4CH2OCH2C4H5	C ₆ H ₆ CH ₂ OC ₁₀ H ₇ - $\beta$ (C ₆ H ₆ ) ₅ COCH ₂ (CHOCOCH ₅ ) ₄ CH ₂ OC(C ₆ H ₆ ) ₅ Note: References 84–165 are listed on pp. 325–326.

* Hydrogenolysis and hydrogenation of the aromatic nucleus are competing reactions. Hydrogenation may follow hydrogenolysis, but not vice versa. t in ethanol no cleavage occurred, and more highly hydrogenated products were formed. I When this product was hydrogenated in ethanol for one hour with Pd black, \$-glyceroglucogide was formed.

TABLE V-Continued

8			ORGANIC	REACT	ONS		
	Refer- ence 15	101	101	101	101	101	101
	Time 32 hr.	2-3 hr.	2-3 hr.	2-3 hr.	2-3 hr.	2-3 br.	2-3 hr.
	Pressure atm.	ı	1	ı	1	1	1
	Temper- ature °C, 40-50	40-50	40-50	40-50	40-20	40-50	40-50
	Solvent CH ₃ CO ₂ H	Ahs, ethanol	Abs. ethanol	Abs. ethanol	Abs. ethanol	Abs. cthanol	Pd-charcoal Abs. ethanol
	Catalyst Pt hlack	Pd-charcoal Abs. ethanol	Pd-charcoal Abs. ethanol	Pd-charcool Abs. ethanol	Pd-charcoal Abs. ethanol	Pd-charcoal Ahs. cthanol	Pd-charcoal
	Yield %	92	8	91	93	82	83
	Benzkl Ekhens  Product Isolated  CH40CH(GH0COCH3)*CHCH20H	CH ₂ OH	CH20COC17436 CH20COC17436 CH20H CH20COC6476	CH20COC6H5 OH2OH    -  -	CH2OCOC17H3s CH2OH   	CH2OCOC18H31 CH2OH CHOCOC6H5	CH20C0C17H155 CH20C0C17H156 CH0CC0C17H166 CH20C0C6H16
	Substace Reduced CII,9QII(QHQQQQCIA)3,QHCH2QQC(GeH6)3	CII ₂ OC(C ₆ H ₈ ) ₃	CH0COC1,H1s CH2OCOC1,H2s CH2OCOC3,H2s	CII_2OCOC_#II_s CII_2OCOC_#II_s CII_2OC(C_#II_s)	CH2OCOC17H3s CH2OC(C6H9)3 CH2OC(C6H9)5	   CH2OCOC1,6H31   CH2OC(C,6H3)3   CHOCOC,6H6	CH20C0C ₁₇ H3s CH20C0C ₁₇ H3s CH20C0C ₁₇ H3s CH20C0C ₁₇ H3s CH20C0C ₂ Hs

## TABLE V-Continued

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STIL
XE.]
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	Rofer- enco 111	11	112	E1	33
	Timo 16 min.	20 min.	ì	ı	35 min.
Pres-	suro atm. 1	-	I	1	က
Tompor- Pres-	oc.	t	ı	l	1
	Solvent CH1CO2H	си,сол	сп,0и	Abs. ethanol	Pd-charval CH4CO2H +
	Yield Gatalyst Solvon % Catalyst Solvon Quant. Pd-charcoal CH4CO2H	Pd-charcoal CH4CO2H	Quant. Pd-charcoal CH4OII	Pd-charcoal Abs. ethanol	Pd-charcoal
	Yield % Quant.	1	Quant.	96	72
Benzyl Ethers	Product Isolated OCH,	110 CO ₂ CO ₂ H ₅	HOCH3	CH ₂ OCH ₃	OCH3, OH
	Substance Reduced	00140000000000000000000000000000000000	N ₂ OCH ₃	In Oction	OCH ₂ C ₆ H ₆

TABLE VI Acetals

					pera-	Pres-		,
Substance Reduced	Product Isolated	Yield %	Catalyst Pd-charcoal	Solvent CH ₃ CO ₂ H	ture C.	sure atm.	Time 30 min.	Keter- ence 33
C,H,CH(OC,H;),CHOHC,H,	2C ₂ H ₅ OH		Pd black Pt black	+ HClO ₄ CH ₃ CO ₂ H CH ₃ CO ₂ H	1 %	1 (	11	10 40
Con Control	C6H11CH3 + C6H11CH2UA + C4H11CH2OC2H6	- · 	o Diach	TOO HO	1	١	ţ	10
$p\text{-}\mathrm{CH}_3\mathrm{C}_6\mathrm{H}_4\mathrm{CH}(\mathrm{OC}_2\mathrm{H}_5)_2 \ p\text{-}\mathrm{CH}_3\mathrm{OC}_6\mathrm{H}_4\mathrm{CH}(\mathrm{OC}_2\mathrm{H}_5)_2$	p-CH ₃ C ₆ H ₄ CH ₃ + 2C ₂ H ₅ OH p-CH ₃ OC ₆ H ₄ CH ₃ + 2C ₂ H ₅ OH	11	Pd black Pd black	CH3CO2H	11	1	1	12
OTHO				•	,		c c	Ę
C,H,CO2CH CHC,Hs	C6H5CO2CH(CH2OH)2	98 Pd	q	Abs. ethanol	25	<b>-</b>	1-z nr.	<b>T</b>
CH20				:	è	-		Ę
CH2O	CH3CO2CH(CH2OH)2	Pd I	77	Abs. ethanol	63	<b>-</b>	l	Į.
CH3CO2CH CHC4H3								
CH20						,		:
OH2	C ₁₅ H ₃₁ CO ₂ CH(CH ₂ OH) ₂	96 P.	Pd black	Abs. ethanol	25	-	90 min.	4
C15H31CO2CH CHC6H5								
CH2O					;	,		ç
Benzal-α-methylglucoside C ₆ H ₅ CH(OCOCH ₃ ) ₂	$lpha ext{-Methylglucoside}  ext{C}_6 ext{H}_5 ext{CH}_3$	<u>r</u>	Pt sponge —	Ethanol —	25		11	7. 7.

Note: References 84-165 are listed on pp. 325-326.

### HYDROGENOLYSIS OF BENZYL GROUPS

## TABLE VII

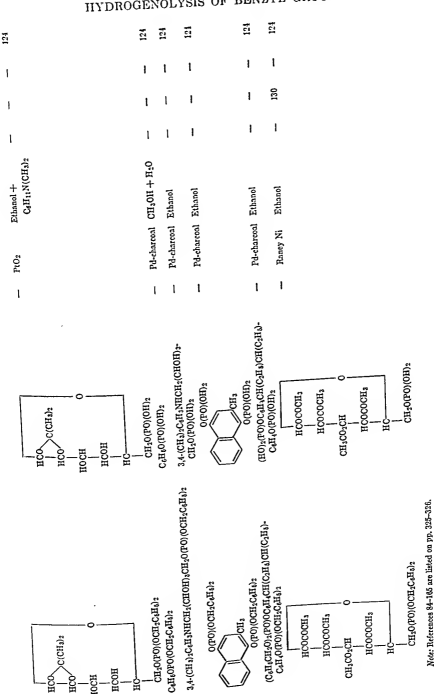
HIDROGENOUTES OF			
Reference 7 9 43 43 43 43 117 118 119 120 120 120	120 120 120	120 120	44, 121
Time	45 min. 24 br. 4 d.	7 d. 3 br.	1
Pres-	1 1 1	18	i
Temper P ature 8 °C. 9 140 150-215 150-215 150 150 150 150 150 150 150 150 150 1	25 25 25	22 22	22
Solvent CH ₃ CO ₂ H Xylene Xylene Xylene Tetralin Etbanol + HCl Dil. CH ₃ OH H ₂ O Abs. etbanol	Pd-cbarcoal Abs. ethanol Pd-charcoal Abs. ethanol Pd-charcoal Abs. ethanol	Pd-charcoal Abs. ethanol Pd-charcoal Abs. ethanol	Pd-charcoal Etbanol + etbyl acetate
Catalyst Pd Pd Pd Pd-BaSO ₄ Pd-BaSO ₄ Pd-BaSO ₄ Pd-BaSO ₄ Pd-Charcoal	Pd-charcoal Pd-charcoal Pd-charcoal	Pd-chare Pd-chare	Pd-char
Yield % CG % CG + Pd + P	97 Quant. Quant.	1-1	91
Benzyl Esters  Product Isolated Cehech, Cehech, Ceheco, Cehecheco, Cehecheco, Product College Cehecheco, Cehecheco, Cehecheco, College, Chico, College, Chico, College, Chico, Cehecheco, C	C ₂₃ H ₃₆ (OH)CO ₂ H (hydroxycholenie acid) C ₂₃ H ₃₆ (OH) ₃ CO ₂ H (cholie acid) C ₂₃ H ₃₆ (OCOCH ₃ ) ₃ CO ₂ H (triacetylcholie	acid) C ₂₈ H ₄₆ CHOHCO ₂ H (oleanolic acid) C ₂₈ H ₄₄ O(CO ₂ H) ₂ (quinovic acid)	n-C,H ₁₅ COC ₉ H ₁₉ -n
Substance Reduced Collocidio Coll	C23H3c(0H)CO2CH(C2H3)2 C23H3c(0H)3CO2CH(C3H3)2 C3-H3-c(OCOCH3)2CO2CH(C6H3)2	C. M. COO-CU(C-Us)	n-C-111, COC(C ₃ 111 ₇ -n)(CO ₂ CU ₂ C ₆ H ₆ ) ₂

Nete: References 84-165 are listed on pp. 325-326.
• Under the same conditions benzaldebyde was reduced via benzyl alcohol to dibenzyl ether.

† The yield is a function of time and temperature.

VII—Continued
TABLE

OR	GANIC R	EACTI	10	IS						
Refer- ence	44, 121 44, 121	44, 121	121	44	44	44	44	122 122	122	
Time	1.1	i	1	i	1	١	1	11	10 min.	1
Pres- sure	1 1	i	ı	1	ı	ı	١	1 1	·	-
Temper- I	22 22	26	25	i za	ĸ	ĸ	32	1 1	1.1	1
tuan lo	thyl	Pd-charcoal Ethanol	i	Pd-charcoal Ethanol + ethyl Pd-charcoal Ethanol + ethyl acetate	PAL harcoal Ethanol + ethyl	Pd-charcoal Ethanol + ethyl	acetate Pd-charcoal Ethanol + ethyl	acetate Pd-charcoal Ethanol	Pd-charcoal CH2OH Pd-charcoal CH2OH	arcoal
Yield	28 82	99		81 78	ç	8 8	3 8		1	ة <b>ا</b>
BENZYL ESTERS	Product Isolated n-C ₁₀ U ₃₁ COCH ₂ CH ₂ CH(CH ₃ )2 n-C ₉ H ₁₉ CO(CH ₃ ) ₅ COC ₉ H ₁₉ -n	n-CloH21COCH2CH2CO2H		C2H6OCO(CH2)8COC6H17-n 2-C.H.CH(C2H6)CO(CH2)10CO2H		HO(CH1)10COC3H14-11	Conformation Contracts	m-CH1,0C6H1COCH2CH2CU2H	C14,2CHCH,CH2O(CO)(OH)(OCH2C,Ha)	C.H.Q(PO)(OH)2 C.H.Q(PO)(OH)2
	Substance Reduced Substance Reduced F.O.16II.11(CHI.1).11(CO.1CI.1).1 F.O.16II.11(CHI.1).11(CO.1CI.1).1	H2C4H8)2	n-CloH11COC(CO2CH2CeHs)1	CH_CO_CHI-CeH1  CH_CO_CHI-CeH1  CH_CO_CHI-CeH1	C111,CH(CH(C111)COC(Cl0112;CO2CH2C4H)(CO2CH2C4H5)2	CH-CO-(CH2)10COC(C4H17-n)(CO2CH2C4H8)2	CAH CH (OCOCH DCOC(C12H26-n)(CO2CH2C4H b)2	m-CH,OCeH ,COC(CO;CH;CeH);CH;CO;CH;CeHs	(C,H,GH,GH,2O);PO(OC;H,A) (C,H,GH,2O);PO(OCH,GH,GH(CH,A);	p-C ₁₀ H ₁ OPO(OCH ₂ C ₆ H ₀ ) ₂ C ₂ H ₄ OPO(OCH ₂ C ₅ H ₀ ) ₂ C ₂ H ₄ OPO(OCH ₂ C ₅ H ₀ ) ₂



# CARBOBENZYLOXY COMPOUNDS

	OF	RGANIC REA	CTIONS			
Refer- enco 125	125	25 th ;	45 45 126 128	127	128 128 128 128	81 81 81
Ra Time c 2-3 d.	1	111	11111	1	30 min. 30 min. 90 min.	11-1
Pres- Bure atm.	1	11-	11111	1	1111	11 1
Tem- pera- 1 ture °C. :	25	118	11111	1	1111	11 1
Solvent	Сизон + сизсози	он+но 1		Сн, он + ист	CH ₂ OH + CH ₃ CO ₂ H CH ₃ OH + CH ₃ CO ₂ H CH ₃ OH + HCl Dil. CH ₃ CO ₂ H	Ethanol + CH ₃ CO ₂ H CH ₃ CO ₂ H —
Catalyst	Pd black Pd black	Pd Pd Pd black	Pd Pd Pd Pd black Pd black	Pd black	Pd black Pd black Pd black Pd black	Pd black Pd black Pd black
TYNDS  Xield  %	1 2	Quant. Quant. Quant.	Quant. Quant.	1 1	95 92 Quant.	90 89 75
CARBOBENZYLOXY COMPOUNDS  Yield Product Isolated %	5=5	Hand CCH;  Hand CCH;  NH CCH;  NH CCH;  NH CCH;  PHOCHCH(NH)O0;  PHOCHCH(NH)O0;  PHOCHCH(NH)O0;  PHOCHCH(NH)O0;  PHOCHCH(NH)O;  PHOCHCH(NH)O;	P-HOCENTACE CONTROLLEGORE  H ₂ NOFI(CONTROLLEGORE  H ₂ NOFI(CONTROLLEGORE)  H ₂ NOFI(CONTROLLEGORE  H ₂ NOFI(CONTROLLEGORE  H ₂ NOFI(CONTROLLEGORE)	H ₂ N(CH ₂ ),CH(CONH ₂ )NHCOCe ⁴¹ s  CH ₂ —CH ₂ H ₂ C(C ₂ —CH ₂ NCOCH ₂ NH ₂	CH2—CH2 H2NCH(CH3)CO3 (CH3)2C(CO2H)N (CH3)2C(CO2H)N	HO2CCH2CHCONHCH2CO2C3H3NH2 HO2C(CH3)2CH(CONHCH2CO2C3H3NH2 H3_CCHCH2CH(CONHCH2CO2C3H3NH2
	Substance Reduced  H ₃ CCCCCH ₃ C ₆ H ₅ CH ₅ CCHNC  C ₆ H ₅ CH ₇ CNNC  CCH ₃	Chichical Composition (Composition Composition Composi	Call Color C	Cententonical Antonical Antonical Antonical Series Cententonical Cententonical Antonical Series Cententonical Antonical Series Cententonical Series Centento	H ₂ C(	(CH3)2C(NHCO2CH2C4H3)CONHCH2CO2CH2C4L6 HO2CCH2CH(CONHCH2CO2C2H3)NHCO2CH2C4L6 HO2CCH2CH(CONHCH2CO2C2H3)NHCO2CH2C4L6 HO2CCH2CHCCH3)2CH(CONHCH2CO2C2H3)NHC- CO2CH3C4HC (CH3)2CHCC4A

			]	HYI	oro	GEN	OLY	SIS	OF	BENZ	YЪ	GRU	UFK	,			
128	129	126	130	5	131	132	133	133	133	ន		134	134	135	3	135	
30 min.	1	1	١	1	١	١	1	١	1	1		1	I	١	١	١	
30	1			1	1	١	1	1	1			I	1	1	1	1	
1	1	1	1	' 1	1	1	1	١	1	ន		1	١	١	١	١	
CH13011 + CH13CO211		CH10H + HCl	1	,	ı	HCI + CII,CO.H	Ethanol + H-SO4	сизон	Ethanol $+ \text{CH}_3\text{CO}_2\text{H}$	Dil. II.SO4		1	CH3OH + HCI	СН,0Н + СН,СО-Н	١	Dil. CH3CO2H	
P.4 Mack		Pd black	Pd black		Pd black	Pd black	Pď black	Pd black	Pd black	Pd black •		Pd black	Pd black	Pd black	Pd black	Pd black	
	: Z	로 	·		Agrant.		35			08		ì	١	92	Quant.	8	
	(CH ₃ ) ₂ CHCH ₂ CH(NH ₂ )CONHCH(CH ₃ )CO ₂ L ₁	H ₂ NCH ₂ CUNHCH(CC ₂ C ₂ C ₂ C ₃ CC ₁ C ₁ C)		H ₂ N(CH ₂ ),CH(CONH ₂ )NHCOCH ₂ NHCOC ₆ n ₃	1)CH2-	H ₂ NC(=NH)NH(CH ₂ ) ₃ CH(CO ₂ H) ^N H- COCH ₂ NH ₂ H ₂ NC(-NH)NH(CH ₂ ) ₂ CH(CO ₂ H)NH-	COCHANIS	H ₂ N(CH ₂ ),CH(NH ₂ )CONNCH ₂ CO ₂ L	CellsCONE(CH2/CH2/CH2/CH2/CH2/CH2/CH2/CH2/CH2/CH2/	CO-H IIC====CCH2CH(CO-H)NHCO(CH2)3NH2.*	He HN N	CH H2NCH2CONHCH(CONH2)CH2C6H4OH-P	"HOC.H.CH.CH(NH.)CONHCH.CO.C.118	-HOC/T-CHOC'HOCHO'H-	CH(CO ₂ H)CH ₂ CO ₂ H  CH(CO ₂ H)CH ₂ CO ₂ H	CH(CO ₂ II)CH ₂ C ₆ H ₄ OH-p p-HOC ₆ II ₄ CH ₂ CII(CO ₂ H)NHCO-	ave even
	$(CH_3)_2CHCH_2CH(NHCO_2CH_3C_6H_6)CONHCH(CH_3). \qquad (CH_3)_2CHCH_3CH(NH_2)CONHCH(CH_3)CO_2H_6$	CO2H C6H3CH2CONHCH2CONHCH(CO2C2H3)CH2CH2-	COLOLARS C6H6CH2OCONH(CH2),CH(CO2CH3)NHCOCH2NH-	CO ₂ CH ₂ C ₆ H ₅ C ₆ H ₅ CH ₂ OCONH(CH ₂ ) ₄ CH(CONH ₂ )NH-	Cocii2NHCOC,Hs CeHcH2OCONHCH(CH2CH2CO2H)CO-	NHCH(CO2H)CH2CH2CO2H O3NNHC(=NH)NH(CH2)3CH(CO2H)NHCOCH2- NHCO3CH2C6H8	O2NC(=NI)NH(CH2)2CH(CO2H)NHCOCH2- NHCO2CH2C6H8	C ₆ H ₆ CH ₂ OCONH(CH ₂ )4CH(NHCO ₂ CH ₂ C ₆ H ₄ )CO- NHCH ₆ CΩ ₉ H	Conscional (CH2) ACH (NHCO2CH2CoH3) CO-NHCH(CCO2H) CH2CH2CO2H	Consider Strain Constitution of the Constituti	HN N CO2CH2C4H5 HN	CH CH CH . TH TH OLD HICH COONE 2) CH2	Cont. One-p	p-CH ₂ CO ₂ C ₆ H ₄ CH ₂ CH(NHCO ₂ CH ₂ C ₆ H ₅ )CO ₂ NHCH ₂ CO ₂ C ₂ H ₅	P-CH ₂ CO ₂ C ₆ H ₂ CH ₂ CO ₂ CH ₂ C ₆ H ₂ CO ₂ CH ₂ CO	CelliCH20CONHCH(CH2CD2H)CONH- CH(CO2H)CH2CeH4OH-p 2-HOC.H.CH-CH(CO-H)NHCO-	CH(NHCO ₂ CH ₂ CcH ₆ )CH ₂ CH ₂ CO ₂ H  Note: References 84-165 are listed on pp. 325-326.  * Hydrogenolysis with sodium in liquid ammonin gave equal yields of extrosine.

-Continued	7 COMPOUNDS
M	LOX
TABLE	CARBOBENZYLO

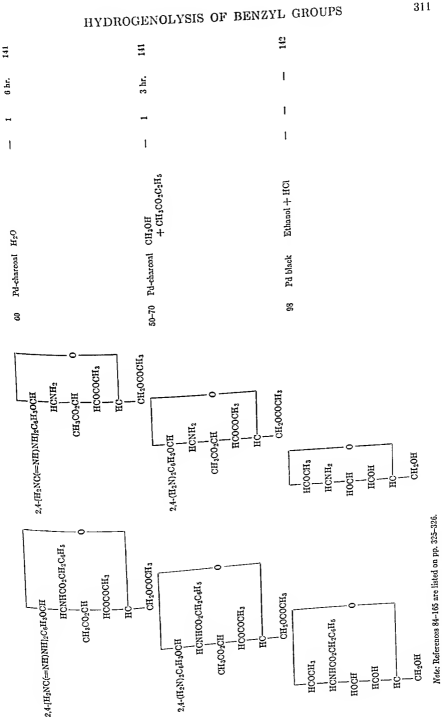
	Refer- ence 47	135	47	18	<b>5</b>	129	18	126	127	47	18	81	129	
	Time R	3-4 d.	1	1	1	ı	1	1	1	l	1	1	1	
Desc	<u>-</u>	بة ا	<b>#</b>	ı	ı	1	ı	1	1	-	1	1	1	
Tem-	pera- rr ture su °C. at 25	1	33		1	ı	ı	1	1	×	1	1	1	
ŭ	od tr Solvent	СН°ОН + СН³СО⁵Н	Ethanol + HCl	Ethanol + CH3CO2H	СН ₂ ОН + СН ₂ СО ₂ Н	HCI	1	Ethanol + HCl	сн,он + исі	Diorans + ethanol + FCl	CH30H + CH3CO2H	СН ₃ ОН + СН ₃ СО ₂ Н	HCI	
	Catalyst Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	Pd black	
UNDS	Yield %	Ount	Quant.	. 75	93	1	1	ı	1	1	1	1	1	
CARBOBENZYLOXY COMPOUNDS	Product Isolated	p-HOC6H4CH2CH(NH2)CO- NHCH(CO2CH3)CH2C6H40H-p	ė.	H- PHOCHICESCH(NH2)CONT- CH(CO2C3H3)CH2C6H4OH-P	-		CH2CO2GH CONHCH2CO2HD CH2CO2GH CONHCHANG		46)NH- H3NCH3NCHCANCELOCATO COCH3NHCOCGH6 NH2	(CH3)2CHCH2CH2CONH- CH(CONHCH2CO2H)CH2CH(CH3)2 CH2C6H4OH-?	)- H2NCHCONHCH(CH2O4H40H-p)- CONHCH(CO2C2H40H2C4H40H-p H2NCHCHCCC2HA0CH1C4C0H1-	CH(CONHCH,CONHCH,CH(CH,1)2 CH(CONHCH,CONH),CH(CONHCH,CO,H)-	CH ₂ CH(CH ₃ ) ₂	H ₂ N(CH ₂ ),CH(CONHCH ₂ );CO- NHCH((CO ₂ C ₂ H ₃ )CH ₂ CH ₂ CO ₂ C ₂ H ₃
	bonde C	P-HOC6H4CH2CH(NHCO2CH2C6H6)CO-	NHCH(CO2CH1)CH2C6H4VHP-P P-CH3CO2C6H4CH2CH(NHCO2CH2C6H8)CONH-	CH(CO2H)CH2CH(OH2) p-CH3CO2CH4CH2CH(NHCO2CH2C6H8)CONH- CH7CO-C-H3)CH3CH4CH3CH3D	HO2CCH2CH(CONHCH2CO2C2Hs)NH- COCH2NHCO2CH2C6Hs	HO2CCH2CH2CH(CONHCH2CO2H)NHCUCH2 NHCO2CH2C6H8	Celechioconechiconechicone- CH(CO.C.He)CH2CC1CO1C1H6	(CH ₂ ) ₂ CHCH(CONHCH ₂ CO ₂ H)NHCOCH ₂ -NHCO ₂ Ch ₂ Co ₂ H ₃	Cochockoni(Ch),CH(CONHCH2CO2CH6)NH- Coch2NHCOCch NHCO2CH5	(CH ₂ ) ₂ CHCH ₂ CHCONHCH(CONHCH ₂ CO ₂ H)- CH ₂ CH(CH ₃ ) ₂ CH ₂ C ₆ H ₄ OCOCH ₆ -p	C ₆ H ₆ CH ₂ OCONHCHCONHCH(CH ₂ C ₆ H ₄ OH-p)- CONHCH(CO ₂ C ₂ H ₆ )CH ₂ C ₆ H ₄ OH-p	Celconeca.coneca.coneca.coneca.coneca.coneca.coneca.coneca.co.en.coneca.co.en.	Conecenter Coneces Con	NHCH(COFO2HCH3\HCOFO2HCS)HCO-COHCH3\HCO-COHC

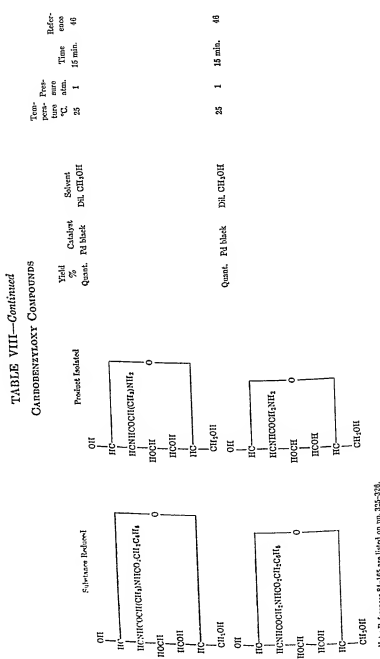
Note: References 84-165 are listed on pp. 325-326.

	ŦŦ	YDROGEN	OLYS	IS OF	BEN	ZYL GRO	UPS		509
47	20 20	136 137 138 138	138	138	139				140
30 min.	1 1	1-6 d. 2 hr. 9 hr. 6 hr.	2 hr. 4 hr.	1 hr.	l				1
E	1 1	- <b>4</b>			i				H
<del>2</del> 2	1 1	ឌ ឌឌឌ	នន	25	1				ĸ
сн ₁ 0н + сн ₁ со ₁ н	Liquid NH3 Liquid NH3		Ethanol + HCl Ethanol + HCl Ethanol + HCl	Ethanol + HCl	i				CH,CO2H + HCI + CH,OH
Pd black	Na Na	PdO Pd-charcoal Pd black	Pd black Pd black Pd black	Pd black	١				Çezak. Pd spozge
ŧ	95 95	89 40 67	8 25 25	<b>i</b>	ſ				il.
NH2   		(CH ₃ ) ₂ CHCH ₂ CH(NH ₃ )CONHCH ₂ - CONHCH ₂ CO ₂ CH ₂ -H ₃ NOH ₂ CO ₂ CH ₃ ₂ -H ₃ NOH ₂ CO ₂ CH ₃		H F	(O-acetyl-r-tyrosyldinydromorphine)  CH2-CH2	Heisconhch(Chichelogy)  Chich(Chis)  Chich(Chis)  Chichich  Cochi	is reduced to CH2—CH2	CHCONECH(CH.C. H.) CHCONECH(CH.): 0 CH—CH:	O-CEL-CECE(NE-CONECH(CO-H)CEL- CE-CE-NE-CONECH(CO-H)CEL- CE-CE-NE-C
NHCO ₂ CH ₂ CeH6	P-CH ₂ CO ₂ C ₂ H ₂ CH ₂ CH ₂ CH ₂ CH ₂ CH ₂ CH ₂	SCH2CH(CO2H)NHCO2CH2CeH5 SCH2CH(CO2H)NHCO2CH2CeH6 CONHCH2CH(NHCO2CH2CeH6)CONHCH2- CONHCH2CO2H CONHCH2CO2H	Cont. CH. COONHCH. CO. Co	Cettly, trist, CONHOLIZ-CONTRACTOR Cettly, to Carbobensyloxgly orlanophine (carbobensyloxgly orlanophine)	CO-C17H18O2N (O-acetylcarbobenzyloxy-r-tyrosylmorphine)	(CH3)*CHCHCONHCH(CH*CH*CH2*OHTCO)*CH4)*CONHCH(CH3*CH4;)CN 	is red	(CH2)-CHCHCONHCH(CH2-CH2-NH2)-CONHCH(CH2-C-H2)-CN	(CH3)-CHCH(NHCO-CH-C ₆ H3)CONHCH(CO-H3)- CH1-CH1-NHCO-CH-C ₆ H3

TABLE VIII—Continued

ę,	enco 140	140	140		140		141						
	Time 1	1	1		1		6-8 hr.						
Pres-	suro stm. 1	-			-		-						
Tem- pera- Pres-	13 C 25	52	22		ន		1						
	Solvent CH3CO ₂ H + HCl	сн,со2н + си,оп	Си,со,п + си,он		сп,со2н + сн,он		сн,0н						
	Catalyst Pd spongs	Pd sponge	Pd sponge		Pd sponge		Quant. Pd-charcoal CH3OH						
UNDS	Yield %	78	: 18		æ		Quant.						
Cahodenzyloxy Compounds	Product Isolated Product Solved		CH(CH ₁ CHCH(Nh ₂ )COM: CH(CH ₁ C ₆ H ₁ )CO ₂ H CH ₂ CH ₂	Caulchichinhalon	C112	CH ₂ CHCONHCH(CO ₂ H)CH(CH ₃ ) ₂	н	p-112NC(==N11)NHC6H40CH	HCNH2	cu _s co _s cu 	II COCOCII:	lio lio	сигососия
	Suleiance Reduced	C4H,CH,CCONH(CH;);CH(NHCO;CH;C4H,)CO-	(CH), CHCH, CH (CH), CONT- CH(CH), CALL) CO, H	Contentantos chicanon Contenta	CO211	CH1 CHCONHCH(CO3H)CH(CH3)2	CO,CII,Calis	P-II,NC(=NII)NIIC,II,OÇII	IIÇNIICOZCII3G4II4	o norootuo	1100000113	) III	 cn2000011





Note: References 81-165 are listed on pp. 325-326.

## TABLE IX

# Monodebenzylation to Primary Amines

Pres-suro

pera-turo °C. Tcm-

1111111111		_	28
6000 7 13 113 113 113 51 53	55 56 56 56 143 144	57	
<u> </u>	11 115	36 hr.	16 hr.
	E       E E	20	<b>**</b>
8	81 1188	20-35	ĸ
Solvent CH3CO2H Ethanol CH3CO2H Ethanol Ethanol CH3CO2H Neutral	Ethanol Ethanol	$CH_3CO_2H$ + $HCI$	сн,0н
Catalyst H2PtCls Pd-charcoal PdO Pd-charcoal Pd-charcoal Pd-charcoal Pd-charcoal	Pd-Pt-charcoal Pd Pd Pd Pd-charcoal PtO ₂	Pd-charcoal	Pd-charcoal
Yield % % % % % % % % % % % % % % % % % % %	Quant.	06	1
Product Isolated C ₆ H ₁ CH ₃ + C ₆ H ₁ NH ₂ + C ₆ H ₁ 1NH ₂ + C ₆ H ₁ CH ₃ + C ₆ H ₆ NH ₂ C ₆ H ₆ CH ₃ + C ₆ H ₆ NH ₂ 3,4-(CH ₂ O ₂ )C ₆ H ₃ CH ₃ + C ₆ H ₆ NH ₂ a _c C ₁ C ₁ H ₇ NH ₂ + C ₆ H ₆ CH ₃ H ₂ N(CH ₂ ) ₂ CH(CH ₃ )N(CH ₃ ) CH ₂ CH ₂ CH ₃ NH ₂ CH ₂ CH ₃	H H2NCH2CH2CO3C2H5 CH3CH(NH2)CH2OH C2H3CH(NH2)CH2OH (CH3)2CHCH2CH(NH2)CH2OH H2NCH(CH2OH)CO2H NNT2 ONH2	H ₂ NCH(CO ₂ H)CH(CO ₂ H)NH ₂	CH3(CH2)3CH(NH2)CH2OH
Substance Reduced Cohis MHCH-2Cohis Cohis WHCH-2Cohis Cohis WHCH-2Cohis Cohis WHCH-2Cohis A;4-(CH2O2)Cohis Chis Cohis WHCH-2Chis Cohis WHCH-2Cohis Cohis WHCH-2Cohis Cohis WHCH-2Cohis	H  H  C ₆ H ₆ CH ₂ NHCH ₂ CH ₂ CO ₂ C ₂ H ₅ CH ₃ CH(CH ₂ OH)NHCH ₂ C ₆ H ₅ C ₅ H ₅ CH(CH ₂ OH)NHCH ₂ C ₆ H ₅ C ₅ H ₅ CH ₂ CHCH(CH ₂ OH)NHCH ₂ C ₆ H ₅ C ₆ H ₅ CH ₂ NHCH(CH ₂ OH)CO ₂ H  NHCH ₂ C ₆ H ₅ NHCH ₂ C ₆ H ₅	C ₆ H ₅ CH ₂ NHCH(CO ₂ H)CH(CO ₂ H)NHCH ₂ C ₆ H ₅	CH3(CH2)3CH(NHCH2C4B3)CH2OH

Note: References 84-165 are listed on pp. 325-326.

IX—Continued
TABLE

1		ORGANI	C REA	ACTIONS	
;	Reference enco 145	145		Reference 51 53 53 53 53 51 51 11 11 146	
	Timo 3 hr.	25 min.		Time 6 hr.	
Pres-	atm.	1		Pres-	
Tem-	.c. 60	20		Tem- ture °C. 25 25 25 25 25 25 25 25	
	Solvent CH3CO2H	СИ3СО2И		Solvent Ethanol  Ethanol  CH3CO2H CH4CO2H H2O H2O	
AMINES	Catalyst Pd-charcoal	Pd-charcoal		Catalyst PdO  Pd   Pd   Pd-charcoal PdO	
MARY .	Yield % 90	1		nr Amn 66 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	group.
MONODEBENZYLATION TO PRIMARY AMINES	Product Isolated CH2CONHCH2C6H6	CHCO ₂ H NH ₂ CH ₂ —C=0 CH ₂ —C=0 CH—C=0	$^{ m hH_2}$ TABLE X	DIDEBENZYLATION TO PRIMAI  Product Isolated  Ranch Cohi, Correction To Primai  Cohi, Correction To Primai  Cohi, Correction To Primai  Cohi, Correction To Primai  Hanch Charles Nat  Hanch Charles Nat  Hanch Charles Nat  Hanch Cohi, Cohi  Cohi, Correction  Ranch Cohi, Charles  Ranch Cohi, Cohi  Ranch	25-326.
	Substanco Reduced	CHCO2H NHCH2C6H6 CH2-C=O CH1-C=O CH-C=O	 NHCH4Ce. Note: References 84–165 are listed on pp. 325–326.	Substance Reduced (GeH5CH2)2NCN (OeH5CH2)2NCH2CH4OC6H5 (GeH5CH2)2NCH2CH5OC6H5 (GeH5CH2)2NCH2CH5OCH2NCH4)CH2CH2OH (GeH5CH2)2NCH4CH2CH2CH2NCH3)CH2CH2 (GeH5CH2)2NCH2CH2CH2CH2CH2CH2CH3)2 (GeH5CH2)2NCH2CH2CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3CH3C	p-HOC ₆ H ₄ COCH ₂ N(CH ₂ C, H ₆ ) ² . HOI Note: References 84–165 are listed on pp. 325–326.

Note: References 84-165 are listed on pp. 323-320. * With a higher ratio of catalyst to amine the carbonyl group was reduced to a bydroxyl group.

## TABLE M

1	The second second	mp 4e	<b>5</b>	#* u",
	* ! !	į	\$ \$	I
		g e	1	è
and the state of	i y i i	1	ţ	1
	roots.	ı	1	Lihand
ertiary Aus	Catalyst Pd Pd-Aareed	Pd-2, we of	*	P.10
EXZYL T	P (* 1 1	1	ſ	12
Monodebenzylation of Direnzyl Tertiany Amines	Product Polated Call,CH;NHCH;CH;COCH3	C CHOHOH-MICH-CAI.	Cuicuismentent	N II III III III III III III III III II
	Substance Reduced (GeH,GH,)NCH,CH,CH,COCH,	CHOHOH:NGH;CdH,)2	COCH,N(CH,C,H,);	CH1 N H H CH2 NCH2C6H1

Note: References 84-165 are listed on pp. 325-326.

TABLE XII

# COMPETITIVE DEBENZYLATIONS

	URG.	ANI	L J	LLH	LOI	10	NO								-	
	Refer- ence	16	16	2 92	16	9 9	16	Pi	16	16	16	16	2	59, 147	59, 147	
	Time	1 1	1	1 1	1	1 1	1	ı	1	1	1	ı	1	Slow t	Slow †	
	Pres- sure atm.	<b>63</b> (		n n	63	<b>~</b>	° 60	က	က	က	က	e5 -	က	es 6	90	
Tem-	ture C.	22	8 8	8 %	33	22	3 3	22	75	22	22	22	22	8	8 23	
	Solvent	СН3ОН	CH ₃ OH Ethanol	Ethanol Ethanol	Ethanol	Ethanol	Ethanol + HCl Ethanol	Ethanol	Ethanol	СН3ОН	СН,ОН	CH ₃ OH	снзон	Ethanol	Abs. ethanol CH ₃ 0H	
70		Catalyst Pd-charcoal	Pd-charcoal Pd-charcoal	Pd-charcoal	100-10-10-10-10-10-10-10-10-10-10-10-10-	Pd-charcoal	Pd-charcoal	Pd-charcoal	Pd.charenal	Pd-charcoal	Tod showned	Pd-charcoal	Pd-charcoal	Pd-charcoal	Pd-charcoal Pd-charcoal	
ATIONS	Yield	% Ouant.	l g	1	l	Quant.	Quant.	1 1	1	1 1		Quant.	Quant.	1	30	8
COMPETITIVE DEBENZYLATIONS		Product Isolated	P-CH ₂ C ₆ H ₄ CH ₂ NHCH ₃ ·HCl P-ClC ₆ H ₄ CH ₂ NHCH ₃ ·HCl	P-CH ₃ OC ₆ H ₄ CH ₂ NH ₂ ··········· P-CH ₃ OC ₆ H ₄ CH ₂ NHCH ₃ ····································	m- and p-CH10C6H4CH2NHCH1-HCl	o- and p-CH ₂ OC ₆ H ₄ CH ₂ NHCH ₂ HCl	3,4-(CH102)CH10CH2CH2CH2 p-CH10CcH4CH2NH2-HC1+p-HOCcH4CH3	P-CH ₃ OC ₆ H ₄ CH ₂ NH ₂ ·HCl + p-HOC ₆ H ₄ CH ₃ H NC H CH ₂ NHCH ₁ ·2HCl •	P-H2N Centions and an area	P-CI(CH1)3NG4CH2NH2·HCI	+ p-cen centensulting	Center House	Collocation Colloc	DH'HOHN HO HO HO		)) p-Chatchatchatchar p-Chatchatcharnerared p-Chatchatcharnerared
		Substance Reduced	PCH3CeH4CH2N(CH3)CH2CeH4.HCl	P-CICCHACHENCHING AND	p-CH10C6H4CH2N(CH1)CH2C6H7TOC m-CH10C6H4CH2N(CH3)CH2C6H40CH1-p-HCl	CH OCALLCHAN(CHA)CHACALLOCH PP-HCI	3,4 (CHrO2)CeH2CH2NHCH2CeH4OCH+P·HCl	P-CH10C6H1CH2NHCH2C6H10H-P-HCI	P-02NC6H,CH2N(CH1)CH2C6H6·HCI	p-CI(CH ₃ ) ₂ NC ₅ H ₄ CH ₂ NHCH ₂ C ₆ H ₅ ·HCl	[a-CloH1CH2N(CH3)CH2CentCene-pj: HC1	A-C10H,CH2N(CH3)CH2C6H6·HC1	B-C10H,CH2N(CH3)CH2C6H6-HCI	a-C ₁₀ H ₁ CH ₂ N (CH ₃ )CH ₂ C ₁₀ H ₇ -B·H·C ₁	$_{p}$ -CH ₂ OC ₆ H ₄ CH ₂ N(CH ₃ )CH ₂ C ₆ H ₄ Cl- $p$ -HCl $_{p}$ -CH ₃ OC ₆ H ₄ CH ₂ N(CH ₃ )2CH ₂ C ₆ H ₄ Cl- $p$ -PlCl $_{p}$ -CH ₂ OC ₆ H ₄ CH ₂ N(CH ₃ )2CH ₂ C ₆ H ₄ Cl- $p$ -PlCl $_{p}$ -CH ₂ OC ₆ H ₄ Cl- $p$ -PlCl $_{p}$ -CH ₄ - $p$ -PlCl $_{p}$ -CH ₄ - $p$ -PlCl $_{p}$ -CH ₄ - $p$ -PlCl $_{p}$ -PlCl $_{$	p-CH30CtH4CH3N(CH3)CH2CtH4CO2CH3-p-HCl

			HY	DRO	GEN	OLY	PIP	OF
59, 147	59, 147	59, 147	59, 147	59, 147 59, 147	59, 147	59, 147	59, 147	
Fast †	Modernto † 59, 147	Fast †	Moderate †	Slow † Slow †	Slow †	Slow †	Slow t	
89	က	es	က	m m	es	es	က	
52	25	65	33	22 22	83	33	22	
Снзон	СН3ОН	Pd-chareoal CH3OH + 10	eq. HCl CH3OH + HCl	-	CH ₃ OH	04.044.90	Pd-chargon CH3OH 22 eq. HCl	
Pd-charcoal	Pd-charcoal	Pd-chareoal	Pd-chareoal	Pd-charcoal	Pd-charecal Pd-eharccal		Pd-chareout	rd-enarcoar
40	8 6 6	6 1	l &	90 85–90	70 10 95-100	65-70	15	8 8
And the second	1 P-CH;CONHCad,CH; P-CH;CONHCad,CH;N(CH;)2·HC! P-CH;OCad,CH;N(CH;)2·HC!	p-CH ₃ OC ₆ H ₄ CH ₂ NHCH ₃ ·HCI	[p-CH30C6H4CH2N(CH3)CH2C6H4NHz-pjzhC1	p-CH ₃ C ₆ H ₄ NH ₂ ·HCl p-CH ₃ OC ₆ H ₄ CH ₂ N(CH ₃ ) ₂ ·HCl	p-Cicht-Channeriann p-Chaoloccht-Channeriann p-Chaoloccht-Cha	P-CH3C6H4NH2·HCl P-CH3C6H4CH2NHCH3·HCl	$_{p}$ - $_{1}$ N $_{6}$ H $_{4}$ CH $_{2}$ NHCH $_{3}$ - $_{2}$ HCI	C6H6CH2NHCH3·HCl p-CH3C6H1NH2·HCl
	P-CH10CoH4CH2N(CH1)2CH1CoH4NHCOCH1-p CH P-CH1CONHCoH4CH2N   P-CH10COH4CH3N(CH1   P-CH10CH4CH2N(CH1	$p ext{-} ext{CH}_10 ext{C}_6 ext{H}_4 ext{CH}_2 ext{N}( ext{CH}_3) ext{CH}_2 ext{C}_6 ext{H}_4 ext{N}0_2 ext{-}p ext{-} ext{HCl}$	$p ext{-} ext{CH}_3 ext{OC}_6 ext{H}_4 ext{CH}_2 ext{N}( ext{CH}_3) ext{CH}_2 ext{C}_6 ext{H}_4 ext{NO}_2 ext{-}p$	[p-CH4OC6H4CH2N(CH3)2CH2C6H4NH3-p]Cl2	p-CH3C6H4CH2N(CH3)CH2C6H4CH-p-HCl p-CH3C6H4CH2N(CH3)CH2C6H4CO2CH3-p-HCl	p-CH3C6H4CH2N(CH3)CH4C6H4NO2-p-HCl	p-CII 3C4U (CH2) (CH2) CH2C4H 1NO2-p-HCI	p-0,NC,H,CH,N(CH,)CH,C,H,Cl-p HCl

be the smounts of critalyst and substrate. Rarely is the rate of reduction a straight line function of timo. For the examples cited in the above table, Baltaly 147 considers that during an early † Although the approximate times for reduction are included where known, the value of this information is only relative for the total time is a function of many factors, among which must stage of debenzylation an absorption of 1 mmole/5 min. or less is slow; 1 mmole/3 min. to 1 mmole/1 min. is moderate; and any absorption taking place more rapidly is fast. • When 13 moles of hydrogen chloride was present, the henzyl group was not removed; when 34 moles of hydrogen chloride was present, the benzyl group was removed.

# MONODEBERZYLATION TO SECONDARY AMINES

į	cuca 01	10	22 22 22	52	: 23 :	25 25	25 25	99	51 6	ដ ដ	3	148
	Time 4 br.	4 hr.	8 hr. 8 hr.	8 hr.	8 hr.	8 hr. 8 hr.	8 hr.	6 hr.	e li	1 1	I	1 1
Pres-	sure atm. I70	177	000	,	- 2	es es	89 89	۱ ۹	1 1	1	l	٦ ١
Tem-	ture °C. 165	105	65-75	65-76	1 -39	65-75 65-75	65-75	0.2	R 1	25 25	1	25.
	Solvent Dioxane	Dioxano	CH3CO1H CH3CO1H	CH3CO2H	CH,CO,H CH,CO,H	CH,CO2H CH,CO2H	CH2CO2H	CH3CO2H CH3CO2H	CH,CO2H CH,CO2H	Ethanol Ethanol	1	Ethanol —
ty Amines	Catalyst Copper chromium	oxido Copper chromium	oxida PtO ₂ PtO ₂	PtO ₂	PdO	PtO ₂	Pt02	PtO ₂ PtO ₃	Pt02 Pt02	Pd-charcoal Pd-charcoal	I	Pt02
CONDAF	Yield		1 1	1 (	8	111	1	Omat	Quant.	Quant. Quant.	1	Quant.
Monoberenzylation to Secondary Amines	Froduct Folated	2,5.Dinethylhydroquinono 7 (OLS)2	Z.J.o. Iffinetory present in the control of the con	CHINIOSHIT-1	CII,NIIC12U16-n CII,NIIC16II13-n	C416NHC4H2-n C416NHC4H2-n	OIDSMINGTH.	n-C(H ₀ NHC ₆ H ₁₁ -n	(n-Cellis)2N11 (n-Cellis)2NH - Cellis(n-Cellis)2	Continued to	CH ₂ CH ₂ NHCH ₃	C ₆ II,CHO 2,5-(CH ₃ C
	Salutane Reduced	2,5-1114(fimethylaminomethyl)hydroquinons	2.Directhylaminemethyl 3,5-dimethylphenol	C4II,CII,N(CII,)C4II-n C4II,CII,N(CII,)C4II1-n	Colfoll, N(Clf.) Chillists	C, II, C, II, N, C, II, I) C, III, n C, II, C, II, N, C, II, S, III, n C, II, C, II, N, C, II, S, II, n	CellicitaN(Calli)Callitan	Cell_Cll_N(C_1ll_+n)Cell_+n	CallaClisN(Calfis-n)2 CallaClisN(Calfis-n)2	C4H3CH3N(CH3H35+n)2 C4H4N(CH3)CH3C4H8	Centan(Crins)CHrOsus Centan(CHr)CHr)CHroens CHro	N H Calicholich(Chan(Chanchas 25-(Chao):ceatgloth)(Chanchas

	HYDROGENOLYSIS OF BENZYL GR	OUPS	319
54		51	53 149
1	1212 12 12 14 1 1 1 1 1 1 1 1 1 1 1 1 1	1	1.1
1	6 8 8 1 3 4 6 5 6 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1	1-1
1	80-90 25 25 25 20-100 90-100 1	1	
١	H2O Abs. CH3OH 2 N HCI H2O H2O Dil. ethanol Dil. ethanol CH3CO2H CH3CO2H	Ethanol	NH ₃
Pd-charcoal	Nickei Pt black Pt black Pd-acacia Nickei Ni: Co: Cu, 10: 6: 1 Nickei Niekei Pd-acacia Pd PdO	PdO	Nickel PtO ₂
1	1881	94	11
	CHOCE HOCE AND CHEST OF THE CHEST OF THE CHEST OF CHEST O	CNH2 C=NH HN NH HN=C C=NH	N—H CehgChanhchachachchchgnnh2 2,5-(Chao)2CehsChohchanhcha
(	CHILD CHORD, NCH3) CH2C6H6  P-HOC6, H, COCH2, NCH3, CH2, C6, H6, HC1 3-F-4-HOC6, H3, COCH2, NCH3, CH3, C6, H6, HC1 3-CH4-HOC6, H3, COCH2, NCH3, CH3, C6, H6, HC1 3, 4-HOC6, H3, COCH2, NCH3, CH3, C6, H6, C6, H3, COCH3, NCH3, CH3, CH3, C6, H6, C6, H6, C6, H6, COCH3, NCH3, CH3, C6, H6, HC1 CAH, COCH (CH3, NCH3) CH2, C6, H6 CAH, LOCH (CH3, NCH3) CH2, C6, H6 CAH, LOCH (CH3, NCH3, CH2, C6, H6 CAH, LH2, CN NCH2, CR1 NCH2, CR	CNII ₂ C=NII C _c H ₆ H ₇ CN NCH ₂ C ₆ H ₆ IIN=C C=NII	N—CH ₂ CeH ₅ (C ₆ H ₃ CH ₂ )NCH ₅ CH ₂ COCH ₃ 2.5.(CH ₃ O) ₂ CeH ₃ COCH ₂ N(CH ₃ )CH ₂ CeH ₆

THE RESERVE THE PROPERTY OF TH

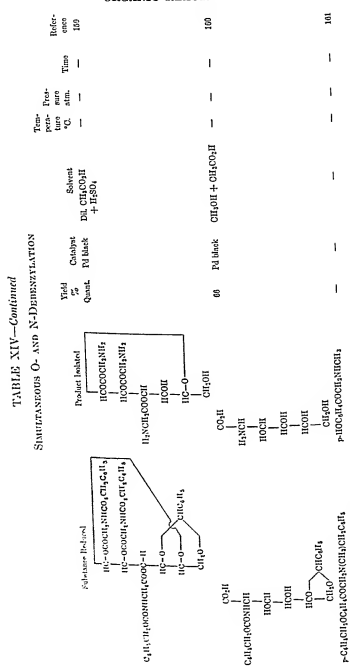
Note: References 84-165 are listed on pp. 325-326.

TABLE XIII—Continued
Monobedentalation to Secondany Amines

Tem-

Refor- ence 54	51	154 155, I56	157	
Time	1.10 min.	1 1	1	
Pres- sure atm.	<b>=</b> =	1 1	1	
rad Sign	1 8	45	1	
Solvent	Ethanol CH3OH	Ethanol Ethanol	СН3ОН	
Catalyst Pd-charcoal	PdO Pd-charcoal	Pd Pd spongo	Pd-charcoal	
Yes	6	87-04 92-98	1	
Project Isolated	(CAIISCII) 1MI	C,U,O;C,U;OCH;NHCH;	H ₈ C ₆ CH ₂ —CH ₂	CII2C6IIs
9 steames Reduced	COLLAND COLLAN	25.(CH,0), CH, CH, CH,  55.(CH,0), CH, CH,  CH, CH, CH, CH,  CH, CH,  CH, CH,  CH, CH,  CH, CH,  CH, CH,  CH,	Hici CH;-CH;	CHICAN

Note: References 84-165 are listed on pp. 325-326.



Note: References 84-165 are listed on pp. 325-326.

TABLE XV
QUATERNARY AMMONIUM COMPOUNDS

Substance Reduced [(C ₆ H ₅ CH ₂ ) ₃ N(CH ₂ )]OH	Product Isolated CeHeCH2NHCH3	Yield % —	Catalyst PdO PdO	Solvent Ethanol Ethanol	Temperature °C.	Pressure atm.	Time — —	Reference
[C6H5CH2N(CH3)2C6H5]Cl	$C_6H_{11}N(CH_3)_2$		Pao	Dillaner				

Note: References 84-165 are listed on pp. 325-326.

### TABLE XVI

## REDUCTIONS WITH NICKEL-ALUMINUM ALLOY 24

REDUCTIONS WITH  Substance Reduced  C ₆ H ₆ CH ₂ OH  C ₆ H ₅ CHO  o-HOC ₆ H ₄ CH ₂ OH  o-HOC ₆ H ₄ CHO  p-HOC ₆ H ₄ CHO  C ₆ H ₆ COCH ₃	Product Isolated  C ₆ H ₆ CH ₃ C ₆ H ₆ CH ₃ o-HOC ₆ H ₄ CH ₃ o-HOC ₆ H ₄ CH ₃ p-HOC ₆ H ₄ CH ₃ C ₆ H ₆ C ₂ H ₅ m-H ₂ NC ₆ H ₄ C ₂ H ₅	Yield % 70 60 85 75 80 70
m-O ₂ NC ₆ H ₄ COCH ₃	TTOC-H.CoHe	72 78
$p ext{-HOC}_6 ext{H}_4 ext{COCH}_3 \ p ext{-HOC}_6 ext{H}_4 ext{COC}_2 ext{H}_6$	$p ext{-}HOC_6H_4CH_2C_2H_5 \ p ext{-}HOC_6H_4CH_2C_6H_6$	90
$p ext{-HOC}_6 ext{H}_4 ext{COC}_6 ext{H}_5$	CH-CH-CH-C6H5	70 50
C ₆ H ₅ COCH ₂ C ₆ H ₅ C ₆ H ₆ CHOHCOC ₆ H ₆	$C_6H_5CH_2CH_2C_6H_6$	50

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    128 D. J. Sahlaish J. Riol. Ch. Bergmann, Zervas, and Fruton, J. Biol. Chem., 109, 325 (1935).

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  - ¹²⁹ Prelog and Wieland, Helv. Chim. Acta, 29, 1128 (1946). 130 Bergmann and Ross, J. Am. Chem. Soc., 58, 1503 (1936).
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#### CHAPTER 6

## THE NITROSATION OF ALIPHATIC CARBON ATOMS

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### Vanderbilt University School of Medicine

#### CONTENTS PAGE 328 NATURE OF THE REACTION 330 330 SCOPE AND LIMITATIONS 336 341 342 343 344 . . . . . . . . . . . . . . . . . . 346 . . . . . . . . . . . . . . . Nitro Compounds . . Hydrocarbons . . . . 346 346 348 349349 352 Dioximinoacetone from Acetonedicarboxylic Acid 352 3-Oximino-5-ethoxy-2-pentanone from Ethyl $\alpha$ -2-Ethoxyethylacetoacetate 353 Ethyl $\alpha$ -Oximinoacetoacetate from Ethyl Acetoacetate . . . 353 354 α-Oximinocaproic Acid from Ethyl n-Butylacetoacetate 354 Ethyl $\alpha$ -Oximinocaproate from Diethyl n-Butylmalonate . . . . . 355 357 358 TABULAR SURVEY 366 Malonic Acids, Esters, and Amides •371 372 Table V. Nitriles 373 Nitro Compounds 374 374 . . . . . . . . . . . . . Table VI. Hydrocarbons . . Table VII.

#### NATURE OF THE REACTION

The nitrosation reaction consists in the replacement of a hydrogen atom by the nitroso group, with the formation of a nitroso or oximino derivative. (Oximes formed by nitrosation reactions have often been called isonitroso compounds. Since isonitroso compounds are identical with oximes produced by other methods, the use of the dual terminology is gradually being discontinued.) With few exceptions, the replacement of hydrogen on an aliphatic carbon atom requires the presence of electron-attracting groups adjacent to the carbon to be nitrosated. Acyl, aroyl, carbonyl, carboxyl, carbalkoxyl, nitro, cyano, imino, and aryl groups may serve as activators, but they vary greatly in their capacity to promote nitrosation. Thus, monoketones are readily converted into α-oximino ketones, whereas monoesters containing no other activating groups do not undergo the reaction.

Victor Meyer discovered the reaction in 1873-1874, when he found that careful acidification of an alkaline solution of a nitroparaffin and an alkali nitrite converts a primary nitroparaffin into a nitrolic acid 1 and a secondary nitroparaffin into a pseudonitrole.2.3 He subsequently

$$\begin{array}{ccc} \text{RCH}_2\text{NO}_2 & \xrightarrow{\text{INO}_2} & \text{RCNO}_2 \\ & & & & & \\ & & & & \text{NOH} \\ \text{R}_2\text{CHNO}_2 & \xrightarrow{\text{INO}_2} & \text{R}_2\text{CNO}_2 \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ \end{array}$$

extended the reaction to  $\beta$ -keto esters by preparing ethyl  $\alpha$ -oximinoacetoacetate from ethyl acetoacetate.4.5

$$\begin{array}{c} \text{CH}_3\text{COCH}_2\text{CO}_2\text{C}_2\text{H}_6 \xrightarrow{\text{HNO}_2} \text{CH}_3\text{COCCO}_2\text{C}_2\text{H}_5 \\ & \parallel \\ & \text{NOII} \end{array}$$

When a methyl or methylene group is nitrosated, the nitroso intermediate usually rearranges rapidly to the oxime. (The isolation of

 $CH_2(CO_2R)_2 \rightarrow ONCH(CO_2R)_2 \rightarrow HON=C(CO_2R)_2$ 

¹ Meyer, Ber., 6, 1492 (1873).

² Meyer and Locher, Ber., 7, 788 (1874). ³ Meyer and Locher, Ber., 7, 1506 (1874).

⁴ Meyer, Ber., 10, 2075 (1877).

⁵ Meyer and Züblin, Ber., 11, 320 (1878).

nitroso intermediates is reported on pp. 333, 338, and 339. The formation of stable nitroso derivatives of two  $\beta$ -diketones is discussed on p. 334.) Formation of an oximino structure frequently occurs even when it necessitates cleavage of the molecule at the carbon which has been nitrosated. Monosubstituted  $\beta$ -keto esters and malonic esters are thus converted into  $\alpha$ -oximino esters. A mechanism for the base-catalyzed

$$\begin{array}{ccccccccccccccl} R & & & R & & \\ R'COCHCO_2R'' & \rightarrow & R'COCCO_2R'' & \rightarrow & RCCO_2R'' \\ & & & & & & \\ NO & & & NOH & \\ & & & & & \\ R'O_2CCHCO_2R' & \rightarrow & R'O_2CCCO_2R' & \rightarrow & RCCO_2R' \\ & & & & & & \\ NO & & & NOH & \\ \end{array}$$

nitrosation and cleavage of a cyclic ketone has been proposed.*,6 That

the cleavage of substituted  $\beta$ -keto esters and malonic esters upon reaction with ethyl nitrite and sodium ethoxide occurs by a similar mechanism is indicated by the isolation of ethyl benzoate and diethyl carbonate after the nitrosation of ethyl  $\alpha$ -benzoylvalerate  7  and diethyl n-butylmalonate, respectively. presumed to be formed from the  $\beta$ -keto ester may be represented by the

^{*}In one of the contributing forms of the resonance hybrid, the nitrogen atom of * In one of the contributing forms of the electrons, thus making possible the electrons organic nitrite is considered to have but six electrons, thus making possible the electrons Woodward and Doering. J. Am. Chem. Soc., 67, 860 (1945). philic attack on the α-carbon atom.

Hauser and Reynolds, J. Am. Chem. Soc., 70, 4250 (1948).

⁸ Shivers and Hauser, J. Am. Chem. Soc., 69, 1264 (1947).

accompanying equation.⁷ The nitrosation of  $\beta$ -keto esters, malonic

$$\begin{array}{c} C_3H_7 & OC_2H_5C_3H_7 \\ C_6H_5COCCO_2C_2H_5 + NaOC_2H_5 \rightarrow C_6H_5C & CCO_2C_2H_5 \rightarrow \\ N=O & ONA^{(+)} \end{array}$$

$$\substack{C_6H_5CO_2C_2H_5+C_3H_7CCO_2C_2H_5\\ N-O^{(-)}N_3^{(+)}}$$

acids, and malonic esters in acid solution has been considered to involve reaction of the nitrosating agent with the enolic forms of these compounds.9-14

Nitrosations have been carried out with nitrous acid, nitrosyl chloride, nitrosylsulfuric acid, nitrous fumes, and esters of nitrous acid. Acid or base is usually added as catalyst with the last two reagents.

#### SCOPE AND LIMITATIONS

Since the principal governing factor in this reaction is the nature of the compound to be nitrosated, rather than the particular reagent used, the following discussion is based upon the types of compounds which undergo the reaction. There has been little study of side reactions; they are discussed briefly in the section on experimental conditions. The conversion of oximino products into the corresponding keto derivatives may be the most significant side reaction, but it is probably not serious if the usual nitrosation procedures are employed.

#### Ketones

A ketone group exerts a strong activating influence in the nitrosation of an adjacent carbon atom. The methylene group of a methyl alkyl ketone is attacked in preference to the methyl group. Diacetyl monoxime, an intermediate in the synthesis of dimethylglyoxime, is prepared in 69-74% yield by the action of ethyl nitrite and concentrated hydrochloric acid on methyl ethyl ketone. 15 (The effects of traces of water

⁹ Barry and Hartung, J. Org. Chem., 12, 460 (1947).

¹⁰ Bouveault and Locquin, Bull. soc. chim. France, [3] 31, 1061 (1904).

¹¹ Meyer and Lenhardt, Ann., 398, 66 (1913).

¹² Onishchenko, J. Gen. Chem. (U.S.S.R.), 11, 197 (1941) [C. A., 35, 7941 (1941)].

¹³ Ritchie, Advances in Enzymol., 7, 95 (1947).

[&]quot;Sidgwick, The Organic Chemistry of Nitrogen, revised by Taylor and Baker, p. 171, Oxford University Press, 1942.

¹⁵ Semon and Damerell, Org. Syntheses, Coll. Vol. 2, 204 (1943).

and of varying the amount of catalyst on the yield of diacetyl monoxime are discussed on p. 351.) When 2,4-dinitrophenylacetone is treated

$$\begin{array}{ccc} \text{CH}_3\text{COCH}_2\text{CH}_3 & \xrightarrow{\text{C}_2\text{H}_5\text{ONO}} & \text{CH}_3\text{COCCH}_3 \\ & & & & & & & & & & & & & & & & \\ \text{CH}_3\text{COCH}_2\text{CH}_3 & \xrightarrow{\text{C}_2\text{H}_5\text{ONO}} & \text{CH}_3\text{COCCH}_3 \\ & & & & & & & & & & & & & & \\ \text{NOH} & & & & & & & & & & & \\ \end{array}$$

with isoamyl nitrite and hydrogen chloride in benzene, an 80% yield of 1-oximino-1-(2,4-dinitrophenyl)-2-propanone (I) is obtained. However, isoamyl nitrite and sodium ethoxide in ethanol lead to the formation of 3-acetyl-6-nitrobenzisoxazole (II) and its decomposition product, 4-nitrosalicylonitrile (III). These compounds also result from the action of sodium ethoxide on the oxime I.

$$\begin{array}{c} \text{CH}_2\text{COCH}_3 \\ \text{O}_2\text{N} & \text{NO}_2 \end{array} \rightarrow \begin{array}{c} \text{CCOCH}_3 \\ \text{O}_2\text{N} & \text{NO}_2 \end{array} \rightarrow \begin{array}{c} \text{COCH}_3 \\ \text{O}_2\text{N} & \text{O}_2\text{N} \end{array} \rightarrow \begin{array}{c} \text{CN} \\ \text{O}_2\text{N} & \text{O}_2\text{N} \end{array}$$

Dialkyl ketones with methylene groups in both  $\alpha$  positions give rise to two isomeric oximino derivatives unless the alkyl groups differ considerably in length or unless one is branched. With alkyl groups of different lengths, nitrosation only of the shorter group is found. 17,18 With alkyl groups of similar size, branching of one of them leads to an oximino derivative formed by nitrosation of the unbranched chain.¹⁷

In a study of ketones containing tertiary carbon atoms adjacent to the carbonyl group, Aston and his co-workers 19,20 found that methyl ketones yield only tertiary nitroso derivatives. Both possible products were isolated from six ketones containing a secondary and a tertiary carbon atom adjacent to the carbonyl group. However, propyl isopropyl ketone and butyl isopropyl ketone underwent only methylenc

Many methyl aryl ketones have been converted into their oximino nitrosation.19 derivatives, but the yields have not always been high. Acetophenols and propiophenols are usually nitrosated in lower yield than are the

Dorsche, Ann., 390, 1 (1912).

17 Ponzio and DeGaspari, J. prakt. Chem., [2] 58, 392 (1898); Gazz. chim. ital., 28, 269 (1898).

¹⁸ Ponzio and DeGaspari, Gazz. chim. ilal., 29, 471 (1899).

¹³ Aston and Mayberry, J. Am. Chem. Soc., 57, 1888 (1935). 20 Aston, Menard, and Mayberry, J. Am. Chem. Soc., 54, 1530 (1932).

corresponding methoxy and halo compounds.21-24 This may be due to ring nitration (probably nitrosation followed by oxidation), since nitrophenols are formed when phenols are allowed to react with amyl nitrite in ether for two or three days.25 Under most conditions acetophenone itself 21,26-32 gives lower yields of oximino derivative than does propiophenone.32-37 Oximinomethyl 4-quinolyl ketone (IV) has been prepared in 60% yield by the action of amyl nitrite and sodium ethoxide on methyl 4-quinolyl ketone.38

A number of substituted phenacyl chlorides have been converted in high yields into the corresponding arylglyoxylohydroxamyl chlorides (V).39,40 Another readily nitrosated group of alkyl aryl ketones is

$$\begin{array}{ccc} \text{ArCOCH}_2\text{Cl} & \xrightarrow{\text{C}_4\text{H}_9\text{ONO}} & \text{ArCOCCl} \\ & & & \parallel \\ & & \text{NOH} \\ & & & \text{V} \end{array}$$

The action of amyl related to 1-indanone (α-hydrindone).40a,41,42,43

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21 Edkins and Linnell, Quart. J. Pharm. Pharmacol., 9, 75 (1936).
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23 Pictet and Gams, Ber., 42, 2947 (1909).

²⁶ Bernton, Arkiv Kemi, Mineral Geol., 7, No. 13, 1 (1918) [C. A., 14, 2168 (1920)].

²⁷ Claisen, Ber., 20, 252 (1887).

28 Claisen, Ber., 20, 656 (1887).

²⁹ Claisen, Ber., 38, 696 (1905).

30 Claisen and Manasse, Ber., 20, 2194 (1887).

31 Hartung, Munch, Deckert, and Crossley, J. Am. Chem. Soc., 52, 3317 (1930).

32 Slater, J. Chem. Soc., 117, 587 (1920). 33 Behr-Bregowski, Ber., 30, 1515 (1897).

34 Claisen and Manasse, Ber., 22, 526 (1889).

35 Edkins and Linnell, Quart. J. Pharm. Pharmacol., 9, 203 (1936).

35 Hartung and Crossley, Org. Syntheses, Coll. Vol. 2, 363 (1943).

37 Hartung and Munch, J. Am. Chem. Soc., 51, 2262 (1929).

33 Rabe and Pasternack, Ber., 46, 1031 (1913).

39 Levin and Hartung, J. Org. Chem., 7, 408 (1942). 40 Levin and Hartung, Org. Syntheses, 24, 25 (1944).

40a Kipping, J. Chem. Soc., 65, 492 (1894).

41 Braun and Kirschbaum, Ber., 46, 3045 (1913).

42 Gabriel and Stelzner, Ber., 29, 2604 (1896).

43 Perkin and Robinson, J. Chem. Soc., 91, 1073 (1907).

²² Hartung, Munch, Miller, and Crossley, J. Am. Chem. Soc., 53, 4149 (1931).

²⁴ Zenitz and Hartung, J. Org. Chem., 11, 444 (1946). ²⁵ Ajello and Sigillò, Gazz. chim. ital., 69, 65 (1939).

nitrite and hydrochloric acid on 5,6-dimethoxy-1-indanone leads to the oximino derivative (VI) in almost quantitative yield.43

no derivative (VI) in almost quasi-
$$\begin{array}{c} \text{CH}_2\\ \text{CH}_3\text{O} \\ \text{CH}_2 \end{array} \xrightarrow{\text{C}_5\text{H}_{11}\text{ONO}} \begin{array}{c} \text{CH}_3\text{O} \\ \text{CH}_3\text{O} \end{array} \xrightarrow{\text{CO}} \begin{array}{c} \text{CH}_2\\ \text{CO} \\ \text{VI} \end{array}$$

 $\beta$ -Diketones usually give good yields of oximino derivatives.⁴⁴⁻⁵² Nitroso intermediates (isolated as the dimers unless otherwise noted) may be obtained if the diketones in ether solution are treated with nitrous fumes.⁵⁰ Nitrosodibenzoylmethane (VII) has been prepared in this manner in 50-60% yield. Alkali, ammonia, or boiling ethanol converts this product into the corresponding oxime VIII. Further

converts this product into the correction 
$$C_6H_5COCH_2COC_6H_5 \xrightarrow{N_2O_3} C_6H_5COCHCOC_6H_5 \xrightarrow{KOH} C_5H_5COCCOC_5H_5$$
 $C_6H_5COCHCOC_6H_5 \xrightarrow{KOH} C_5H_5COCCOC_5H_5$ 
 $C_6H_5COCHCOC_6H_5 \xrightarrow{KOH} C_5H_5COCCOC_5H_5$ 

treatment of this oxime with nitrous fumes yields diphenyl triketone. This reagent effects, in one step, the quantitative conversion of p-nitrodibenzoylmethane into the corresponding triketone. 50 Methone (IX) has been nitrosated in 99% yield by potassium nitrite and hydrochloric  $CH_3$ acid.45  $CH_3$ 

The nitrosation of 1,3-indanedione to the 2-oxime 53,54 is of interest as a potential route to ninhydrin. Unfortunately, all attempts to hy-

[&]quot;Ceresole, Ber., 17, 814 (1884).

⁵ Haas, J. Chem. Soc., 91, 1437 (1907).

[&]quot; Kuster, Z. physiol. Chem., 155, 157 (1926).

¹⁷ Lifschitz, Ber., 46, 3233 (1913). ** Neurille and Pechmann, Ber., 23, 3378 (1890).

Sachs and Herold, Ber., 40, 2714 (1907).

⁵⁰ Wieland and Bloch, Ber., 37, 1524 (1904). ¹¹ Wolff, Bock, Lorentz, and Trappe, Ann., 325, 134 (1902).

¹² Zanetti, Gazz. chim. ilal., 23, 303 (1893). 1 Teeters and Shriner, J. Am. Chem. Soc., 55, 3026 (1933).

⁴ Wislicenus, Ann., 246, 353 (1888).

drolyze the nitrosation product were unsuccessful.⁵³ This stability towards hydrolysis has been attributed to the presence of a nitroso group rather than an oximino group in the 2 position.⁵⁵ The nitrosation product is oxidized to 2-nitro-1,3-indanedione by nitric acid and even by nitrous acid, which usually converts oximes to ketones. 2-Nitro-1,3-indanedione is reduced to the nitrosation product by formic acid. Another nitroso compound which does not rearrange to the oximino form in aqueous acid is the 4,9-dinitroso derivative obtained in 89% yield by the action of nitrous acid on 3,5,8,10-tetraketo-3,4,5,8,9,10-hexahydropyrene.⁵⁶

Cyclic ketones appear to be preferentially nitrosated at a tertiary carbon atom. Baeyer converted menthone into nitrosomenthone (X) in 40% yield by means of ethyl nitrite and acetyl chloride  57  and into  $\beta,\zeta$ -dimethyl- $\epsilon$ -oximinocaprylic acid (XI) in 60% yield by means of ethyl nitrite and hydrochloric acid. However, other workers have

$$(CH_3)_2CH \longrightarrow (CH_3)_2CHCCH_2CH_2CHCH_2CO_2H \longrightarrow (CH_3)_2CH \longrightarrow (CH_3)_2CH \longrightarrow (CH_3)_2CHCCH_2CHCH_2CO_2H \longrightarrow (CH_3)_2CH \longrightarrow (CH_3)_2CH \longrightarrow (CH_3)_2CHCH_2CO_2H \longrightarrow (CH_3)_2CH \longrightarrow (CH_3)_2CHCH_2CO_2H \longrightarrow (CH_3)_2CH_2CO_2H \longrightarrow (CH_3)_2CO_2H \longrightarrow (CH_3)_2CO_2H \longrightarrow (CH_3)_2CO_2H \longrightarrow (CH_3)_2CO_2H \longrightarrow (CH_3)_2CO_2H \longrightarrow (CH_2)_2CO_2H \longrightarrow (CH_$$

reported that the conversion of menthone into this oximino acid is poorly effected by amyl nitrite and hydrogen chloride but is accomplished in 68% yield by amyl nitrite and sodium ethoxide. The nitrosation of pulegone (XII) is interesting in that it yields a derivative of isopulegone (XIII). The base-catalyzed isomerization of pulegone to isopulegone apparently is sufficiently rapid for nitrosation to occur at the

$$\begin{array}{c} CH_{3} \\ CH_{3} \\ CCCH_{3} \\ CH_{3} \\ CCCH_{3} \\ CH_{3} \\ CH_{2} \\ CH_{3} \\ CH_{2} \\ CH_{3} \\ CH_{4} \\ CH_{5} \\ C$$

⁵⁵ Wanag and Lode, Ber., 72, 49 (1939).

Vollmann, Becker, Corell, and Streeck, Ann., 531, 85 (1937).
 Baeyer, Ber., 28, 1586 (1995).

⁵³ Baeyer and Manasse, *Ber.*, 27, 1912 (1894).

⁵⁹ Clarke, Lapworth, and Wechsler, J. Chem. Soc., 93, 30 (1908).

newly formed tertiary carbon rather than at the  $\alpha$ -methylene group.^{59a} N-Acetyl-10-oximinodihydrohomomeroquinene ethyl ester (XV), a key intermediate in the synthesis of quinine, is prepared in 68% yield by the nitrosation of cis-N-acetyl-7-keto-8-methyldecahydroisoquinoline (XIV).6 An exception to the usual nitrosation of the tertiary carbon

of cyclic ketones is observed in the reaction of (-)-epicamphor (XVI), which yields (-)-3-oximinoepicamphor (XVII) on treatment with amyl nitrite and sodamide in ether. 60 However, the tertiary carbon in this ketone is at the bridgehead of a fused ring system.

There have been several reports of the synthesis of  $\alpha,\alpha'$ -dioximino ketones by the nitrosation of monoketones. 61-65 (The synthesis of dioximinoacetone from acetonedicarboxylic acid and the failure to prepare ethyl  $\alpha, \alpha'$  dioximinoacetonedicarboxylate from ethyl acetonedicarboxylate from ethy dicarboxylate are discussed below.) The isolation of  $\alpha, \alpha'$ -dioximinotronic transfer are discussed below.) tropinone (XVIII) from the nitrosation of tropinone was useful in the proof of structure of tropinone, for it indicated that the carbonyl group was located between two methylene groups. 65 The use of amyl nitrite and hydrogen chloride in glacial acetic acid led to this dioxime in 90% yield. The same conditions have been used to convert 2,2,6-trimethyl-

Similarly, treatment of pulcgone with hydroxylamine hydrochloride and excess was Similarly, treatment of pulcgone with hydroxylamine hydrochloride and excess was Similarly, treatment of pulcgone with hydroxylamine hydrochloride and excess was similarly to the state of the state potassium hydroxide yields the oxime of isopulegone. Wallach, Ann., 365, 240 (1909).

⁶⁰ Bredt and Perkin, J. Chem. Soc., 103, 2210 (1913).

Borsche, Wallach Fest., 1909, 301 [Chem. Zentr., 1909, II, 1549].

Harries and Groschuff, Ann., 417, 151 (1919).

Kötz, Nussbaum, and Takens. J. prakt. Chem., [2] 90, 357 (1914).

⁴ Wieland, Ber., 37, 1145 (1904).

[&]amp; Willstätter, Ber., 30, 2698 (1897).

4-piperidone (vinyldiacetonamine) into its dioximino derivative XIX in 60% yield. 62

H₂C—CH—C=NOH HON NOH

NCH₃ C=0

H₂C—CH—C=NOH

$$H_3$$
C

 $H_3$ C

The activating effect of an ammono-ketone (ketimino) group is illustrated by the reaction of 2-methyl-3,3-dimethylpseudoindole (XX) with sodium nitrite and acetic acid. The conversion of 1,3,3-trimethyl-

$$(CH_3)_2 \xrightarrow{HNO_2} CH = NOH$$

$$XX$$

2-methylenedihydroindole (XXI) into the aldoxime XXII in 96% yield ⁶⁷ may be considered as proceeding by way of the quaternary salt XXIII which has the structure of an ammono-ketone.

#### $\beta$ -Keto Acids, Esters, and Related Compounds

The nitrosation of unsubstituted  $\beta$ -keto esters yields  $\alpha$ -oximino- $\beta$ -keto esters, whereas  $\alpha$ -substituted  $\beta$ -keto esters are converted into  $\alpha$ -oximino esters. ^{67a} If the  $\beta$ -keto ester is first hydrolyzed to the  $\beta$ -keto acid,

$$\begin{array}{c} \text{RCOCH}_2\text{CO}_2\text{R}' \to \text{RCOCCO}_2\text{R}' \\ & \parallel \\ \text{NOH} \\ \\ \text{RCOCHCO}_2\text{R}' \to \text{R"CCO}_2\text{R}' \\ & \parallel \\ \text{NOH} \end{array}$$

⁴ Plancher and Bettinelli, Gazz. chim. ital., 29, 113 (1899).

E Kuhn, Winterstein, and Balser, Ber., 63, 3182 (1930).

For The one exception to this generalization is the reaction between unsubstituted nectoractic exters and nitrosylsulfuric acid in sulfuric acid, which leads to oximinoacetic exters in good yield. Bouveault and Wahl, Bull, soc. chim. France, [3] 31, 675 (1904).

treatment with nitrite yields an  $\alpha$ -oximino ketone.^{5,68} This reaction has been developed into a general method for the synthesis of  $\alpha$ -oximino ketones. 69,70 It permits the preparation of 3-oximino-2-pentanone (XXIV) from ethyl  $\alpha$ -ethylacetoacetate in 94% yield. 71  $\alpha$ -Oximino

ketones are obtained from  $\beta$ -keto acids even when there is no substituent in the  $\alpha$  position. Thus, dioximinoacetone (XXV) is prepared in 51% yield by the action of nitrous acid on acetonedicarboxylic acid. 72-75

The nitrosation proceeds very rapidly, evolution of carbon dioxide occurring immediately upon the addition of nitrite. Although 1,2-cyclohexanedione monoxime (XXVI) and its derivatives can be prepared by direct nitrosation of the corresponding monoketones, they are also available from the nitrosation of 2-carbethoxycyclohexanones. 76, 77, 78

$$\begin{array}{c} O \\ & \underbrace{\begin{array}{c} 1. \text{ NaOH, NaNO}_2 \\ \hline 2. \text{ H}_2\text{SO}_4 \end{array}} \\ \text{CO}_2\text{C}_2\text{H}_6 \end{array} \begin{array}{c} O \\ \text{NOH} \\ \text{XXVI} \end{array}$$

It should be noted that the success of this reaction depends on the careful exclusion of air from the reaction mixture during saponification. 76

The few reports dealing with  $\beta$ -imino acids and esters indicate that these compounds resemble  $\beta$ -keto acids and esters in their behavior

^{**}Bouveault and Locquin, Bull. soc. chim. France, [3] 31, 1159 (1904).

To Locquin, Bull. soc. chim. France, [3] 31, 1164 (1904).

⁷² Geissman, Schlatter, and Webb, J. Org. Chem., 11, 737 (1946).

⁷ Koessler and Hanke, J. Am. Chem. Soc., 40, 1717 (1918). Mann and Pope, Proc. Roy. Soc. London, 107A, 84 (1925).

Poplar

⁷⁵ Pechmann and Wehsarg, Ber., 19, 2465 (1886).

Geissman and Schlatter, J. Org. Chem., 11, 11, 12, 1937) [C. A., 31, 4960 (1937)].

7 Jaeger and Bijkerk, Proc. Acad. Sci. Amsterdam. 39, 384 (1930) 17 ⁷⁵ Geissman and Schlatter, J. Org. Chem., 11, 771 (1946).

Dueger and Bijkerk, Proc. Acad. Sci. Amsterdam, 39, 384 (1936) [C. A., 30, 6341]

B Jaeger and van Dijk, Proc. Acad. Sci. Amsterdam, 39, 384 (1936) [C. A., 30, 6341] (1936)].

towards nitrosating agents.^{79,80,81} Ethyl  $\alpha$ -cyano- $\beta$ -imino- $\gamma$ -oximino-butyrate (XXVIII) is produced by the action of nitrous acid on mono-ethyl  $\alpha$ -cyano- $\beta$ -iminoglutarate (XXVII).⁷⁹

Benzoylacetimido ethyl ether (XXIX) is reported to yield its oximino derivative (XXX) when treated with amyl nitrite and hydrogen chloride.²⁶ However, potassium nitrite and sulfuric acid lead to the formation of ethyl α-oximinobenzoylacetate (XXXI).

Schmidt and his co-workers  $^{82-85}$  carried out the nitrosation of  $\alpha$ -monoalkyl  $\beta$ -keto esters with nitrous fumes in the absence of solvent and were able to isolate the intermediate monomeric nitroso esters, which were unstable blue or blue-green oils. On standing several days the nitroso

$$\begin{array}{c} R' \\ | \\ RCOCHCO_2R'' \xrightarrow{N_2O_3} R'CHCO_2R'' \\ | \\ NO \end{array}$$

esters underwent both dimerization and rearrangement to the oxime. A trace of alkali brought about very rapid change to the oxime. With this nitrosation technique, it was found that the ease of cleavage of acyl groups decreased in the order: —CHO, —COCH₃, —COC₆H₅.82

Cyclic  $\beta$ -keto esters are usually cleaved to  $\alpha$ -oximino diesters by nitrosation in the presence of alkali alkoxides. 2-Carbethoxy-4-methyl-

⁷³ Baron, Remfry, and Thorpe, J. Chem. Soc., 85, 1738 (1904).

³⁰ Euler and Euler, Ber., 37, 47 (1904).
³¹ Knorr, Ber., 17, 1635 (1884).

[&]quot;Schmidt and Dieterle, Ann., 377, 30 (1910).

[&]quot;Schmidt and Haid, Ann., 377, 23 (1910).

Schmidt and Widmann, Ber., 42, 495 (1909).
 Schmidt and Widmann, Ber., 42, 1886 (1909).

cyclohexanone is converted into diethyl  $\alpha$ -oximino- $\gamma$ -methyladipate in 25-30% yield by the action of nitrous fumes and sodium ethoxide, but almost twice this yield results from the use of ethyl nitrite and sodium ethoxide.86 With 2-carbethoxycyclopentanone (XXXII), ethyl nitrito and sodium ethoxide lead to a 60% yield of diethyl  $\alpha$ -oximinoadipato (XXXIII), whereas ethyl nitrite and acetyl chloride in the absence of solvent permit the isolation of the cyclic nitroso derivative (XXXIV) in 60-80% yield.⁸⁷ The nitroso intermediate can be cleaved to tho oxime in nearly quantitative yield by the action of sodium ethoxide.

$$C_2H_5O_2CCH_2CH_2CH_2CCO_2C_2H_5 \\ NOH \\ XXXIII \\ CO_2C_2H_5 \\ XXXII \\ CO_2C_2H_5 \\ XXXIII \\ CO_2C_2H_5 \\ C$$

Bouveault and Locquin 10,88-91 employed nitrosylsulfuric acid in concentrated sulfuric acid as a reagent for the conversion of  $\alpha$ -monomikyl β-keto esters into α-oximino esters (65-93% yield). Hamlin and Hartung 92 introduced a convenient modification of this procedure in which n-butyl nitrite and 85% sulfuric acid are used as the rengent combination. α-Oximino-δ-chloro-γ-valerolactone (XXXVI), which is used in the synthesis of hydroxyproline, is prepared from a-nearly). δ-chloro-γ-valerolactone (XXXV) by Bouveault's method (67% yield). 63 The reaction of the lactone XXXV with sodium nitrite and dllute sulfuric acid takes an anomalous course, however, since the oxine acetate XXXVII is obtained (81% yield). 4 α-Oximino-γ-butyroluctone,

E Dieckmann and Groeneveld, Ber., 33, 595 (1900).

Bouveault and Locquin, Compt. rend., 135, 179 (1902).

Bouveault and Locquin, Compt. rena., 200, 110 (13) 31, 1049 (1904).

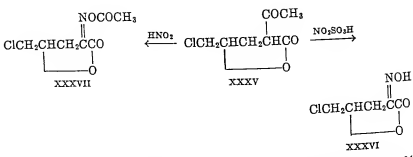
Bouveault and Locquin, Bull. soc. chim. France, [3] 31, 1049 (1904). Bouveault and Locquin, Bull. soc. chim. France, [3] 31, 1055 (1904),

²¹ Locquin, Bull. soc. chim. France, [3] 35, 962 (1906).

²² Hamlin and Hartung, J. Biol. Chem., 145, 349 (1942).

³³ McIlwain and Richardson, Biochem. J., 33, 45 (1939). "McIlwain and Richardson, Biochem. J., 33, 40 (1988) (1988) [C. A., 44] Feofilaktov and Onishchenko, Compt. rend. acad. sci. U.R.S.S., 20, 133 (1988) [C. A., 44]

^{33, 1725 (1939)].} 



an intermediate in a synthesis of methionine, is prepared in 85-91% yield from  $\alpha$ -acetyl- $\gamma$ -butyrolactone, ethyl nitrite, and hydrogen chloride. 95

Diethyl acetonedicarboxylate (XXXVIII) is easily converted to its monoximino derivative by an alkyl nitrite and hydrogen chloride,  96,97  but isoxazole formation occurs when dinitrosation is attempted.  97  The second mole of nitrite obviously serves as an oxidizing agent rather than as a nitrosating agent. The oxime XXXIX and the isoxazole XL have been used in the preparation of  $\beta$ -hydroxyglutamic acid.  96  and  $\beta,\gamma$ -dihydroxyglutamic acid.  98  respectively.

HO_2CCH_2CH—CHCO_2H OH NH2

$$\begin{array}{c} & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & &$$

* Touster and Carter, J. Am. Chem. Soc., 73, 54 (1951).

Snyder, Andreen, Cannon, and Peters, J. Am. Chem. Soc., 64, 2083 (1942).

Harington and Randall, Biochem. J., 25, 1917 (1931).
 Pechmann, Ber., 24, 860 (1891).

 $\alpha$ -Methyltetronic acid (XLI) gives either of two products with nitrous fumes, depending upon the solvent employed.99 A 57% yield of the nitroso derivative XLII is obtained with glacial acetic acid, whereas a 90% yield of  $\alpha$ -oximinopropionylglycolic acid (XLIII) results with water as solvent. It has been reported that other  $\alpha$ -substituted tetronic

livent. It has been reported 
$$\begin{array}{c|c} \text{CO-CH}_2 \\ \text{CO-CH}_2 \\ \text{H}_3\text{CC-CO} \\ \text{H}_3\text{CC-CO} \\ \text{H}_3\text{CC-CO} \\ \text{NO} \\ \text{XLII} \\ \text{XLII} \\ \text{NOH} \\ \text{XLIII} \\ \text{NOH} \\ \text{XLIII} \\ \end{array}$$

acids may suffer loss of the \alpha substituent. 100 Sodium nitrite converted  $\alpha$ -ethyltetronic acid (XLIV) into  $\alpha$ -oximinotetronic acid (65% yield) and acetaldehyde. No explanation was offered for the unusual course of this reaction.

n.
$$\begin{array}{c|c}
CO-CH_2 & CO-CH_2 \\
\hline
O & NaNO_2 & O + CH_3CHO
\end{array}$$

$$\begin{array}{c|c}
CH-CO & C & CO \\
\hline
C_2H_5 & NOH \\
XLIV
\end{array}$$

Malonic Acids, Esters, and Amides

Alkylmalonic acids are decarboxylated during nitrosation. 9, 12, 101, 102

Recent studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have shown that excellent yields of  $\alpha$ -oximino acids can be obtained by the studies have a studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtained by the studies of  $\alpha$ -oximino acids can be obtaine obtained by this reaction. 9,12,101 The action of isopropyl nitrite and hydrogen chloride on 3,4-methylenedioxybenzylmalonic acid furnishes an 85-90% yield of α-oximino-β-(3,4-methylenedioxyphenyl)propionic acid, an intermediate in a synthesis of 3,4-dihydroxyphenylalanine. 101

¹⁰¹ Barry, Mattocks, and Hartung, J. Am. Chem. Soc., 70, 693 (1948).

ter Rietz and Lapworth, J. Chem. Soc., 107, 1254 (1915).

Diethyl oximinomalonate has been prepared from diethyl malonate in good yield under a variety of experimental conditions. Alkyl nitrites, with sodium ethoxide as catalyst, are very effective in converting substituted malonic esters into  $\alpha$ -oximino esters. 9,114 Ethyl  $\alpha$ -oximino-

$$\begin{array}{ccc} \text{RCH}(\text{CO}_2\text{R}')_2 & \xrightarrow{\text{R"ONO}} & \text{RCCO}_2\text{R}' \\ & & \parallel & \text{NOH} \end{array}$$

caproate, ethyl  $\alpha$ -oximino- $\beta$ -phenylpropionate, and ethyl  $\alpha$ -oximino- $\delta$ -dicthylaminovalerate have been prepared in this manner in yields of 80%, 92%, and 94%, respectively.⁸

A number of amides and anilides of malonie acid have been converted into their oximino derivatives. 115, 116, 117 Quantitative yields of oximes were often obtained with nitrosyl chloride as nitrosating agent. 117

#### Arylacetic Acids and Esters

Only a small number of arylacetic acids and esters have been subjected to nitrosation. Ethyl phenylacetate and ethyl p-bromophenylacetate have been converted into their oximino derivatives in good yield by ethyl nitrite and potassium ethoxide.¹¹⁸

$$\begin{array}{c} C_{\mathfrak{e}}H_{\mathfrak{b}}CH_{\mathfrak{c}}CO_{\mathfrak{c}}C_{\mathfrak{c}}H_{\mathfrak{b}} \xrightarrow{C_{\mathfrak{c}}H_{\mathfrak{b}}ONO} C_{\mathfrak{b}}H_{\mathfrak{b}}CCO_{\mathfrak{c}}C_{\mathfrak{c}}H_{\mathfrak{b}} \\ \parallel & \parallel \\ NOH \end{array}$$

Results with nitrophenylacetic acids and esters have not been uniform. Although a few compounds of this type have yielded oximino derivatives

upon treatment with amyl nitrite and a basic or acidic catalyst, 119, 120 others have shown little reactivity towards nitrous acid. 121 usual importance of the nitrosating agent employed is further indicated by the lack of reaction between "2,4-dinitrophenylacetic ester" and isoamyl nitrite and hydrogen chloride in benzene.16 Sodium methoxide catalysis, on the other hand, promotes the conversion of methyl 2,4-dinitrophenylacetate into 3-carbomethoxy-6-nitrobenzisoxazole in 85% yield.16

#### Nitriles

Nitrous acid effects the conversion of methyl and ethyl cyanoacetates into their oximino derivatives in 90% yield, 122-125 but the combination of amyl nitrite and sodium ethoxide leads to poor yields of these products.¹²³ The nitrosation of substituted cyanoacetic esters, like that of

$$NCCH_2CO_2R \xrightarrow{HNO_2} NCCCO_2R$$
 $\parallel$ 
 $NOH$ 

substituted malonic esters, effects decarbalkoxylation, producing the corresponding  $\alpha$ -oximinonitriles. 126 Oximinoarylacetonitriles have been

$$\begin{array}{ccc} \text{RCHCN} & \xrightarrow{\text{R'ONO}} & \text{RCCN} \\ \downarrow & & \downarrow & & \parallel \\ \text{CO}_2\text{C}_2\text{H}_6 & & \text{NOH} \end{array}$$

prepared directly by the action of alkyl nitrites and sodium ethoxide on arylacetonitriles. 127, 128 Nitrous acid converts cyanoacetamides into

The synthesis of oximinomalononitrile (XLV) has been attempted by their oximino derivatives. 122, 129 the nitrosation of malononitrile. Amyl nitrite and sodium ethoxide gave a high yield of a compound assigned the structure  $\alpha$ -oximino- $\beta$ -hydroxy-

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113 Borsche, Ber., 42, 3596 (1909).
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¹²⁰ Gabriel and Meyer, Ber., 14, 823 (1881). Parkes and Aldis, J. Chem. Soc., 1938, 1841.

¹²² Conrad and Schulze, Ber., 42, 735 (1909).

¹²³ Muller, Ann. chim. phys., [7] 1, 463 (1894).

¹²⁵ Fields, Walz, and Rothchild, J. Am. Chem. Soc., 73, 1000 (1951).

¹²⁵ Walker, J. Chem. Soc., 125, 1622 (1924).

¹²³ Zimmermann, J. prakt. Chem., [2] 66, 353 (1902).

¹²³ Merck, Ger. pat. 227,390 [Bril. C. A., 100(i), 166 (1911)].

and addicous access asset

$$\begin{array}{c|c} \text{OH} & \downarrow & \\ \downarrow & \downarrow & \\ \text{C}_2\text{H}_5\text{OC} & \xrightarrow{\text{CCN}} & \xrightarrow{\text{C}_5\text{H}_{11}\text{ONO}} & \text{CH}_2(\text{CN})_2 & \xrightarrow{\text{HNO}_2} & \text{HON} \\ \downarrow & \downarrow & \downarrow & \\ \text{NH}_2 & \text{NOH} & \\ \text{XLVI} & & \text{XLV} \end{array}$$

 $\beta$ -Iminopropionitriles have been found to react with nitrosating agents. ^{132,133} Amyl nitrite in ether converts  $\beta$ -imino- $\beta$ -phenylpropionitrile (benzoacetodinitrile, XLVII) into the ammonium salt of  $\alpha$ -oximino- $\beta$ -nitrosimino- $\beta$ -phenylpropionitrile (XLVIII). *, ¹³² The ammonia necessary for the formation of this compound undoubtedly comes from decomposition of the original nitrile, since oximinobenzoylacetonitrile (XLIX) can also be isolated.

Only a small amount of dioximinosuccinonitrile is formed by the action of two equivalents of amyl nitrite and potassium ethoxide on succinonitrile.¹¹⁹

#### Nitro Compounds

Nitrosation converts primary nitroparaffins into nitrolic acids 1,134 and secondary nitroparaffins into pseudonitroles. 2,3,134 These reactions are the basis of Meyer's "red, white, and blue" test for nitro compounds. 135 Alkaline solutions of nitrolic acids are blood-red in color, whereas pseudonitroles give the blue solutions expected of nitroso compounds. Tertiary nitroparaffins do not undergo nitrosation. Ethyl nitrolic acid (L) 136 (acetonitrolic acid) and butyl pseudonitrole (LI) 3 have been prepared in 82% and 78% yield, respectively. The reaction is carried

¹³⁰ Diels and Borgwardt, Ber., 54, 1334 (1921).

¹³¹ Longo, Gazz. chim. ital., 61, 578 (1931).

¹³² Lublin, Ber., 37, 3467 (1904).

¹³³ Meyer, J. prakt. Chem., [2] 52, 108 (1895).

^{*} A similar compound is reported to be one of the products formed from amyl nitrite and ethyl  $\beta$ -aminocrotonate (ethyl  $\beta$ -iminobutyrate) (see ref. 80).

¹³⁴ Meyer and Locher, Ber., 7, 670 (1874).

¹³⁵ Meyer and Locher, Ber., 7, 1510 (1874).

¹³⁶ Wieland, Ann., 353, 82 (1907).

out by the addition of potassium nitrite and dilute sulfuric acid to an alkaline solution of the nitro compound. When 1,3-dihydroxy-2-nitropropane (LII) is treated in this manner, hydroxyethyl nitrolic acid (LIII) and formaldehyde are formed. 137 The cleavage of the hydroxymethyl group may be similar to that which occurs when other tertiary nitroso intermediates undergo cleavage with rearrangement to the oximes, or it may result from an alkali-catalyzed retrograde aldol condensation prior to nitrosation.

$$\begin{array}{c} \text{NO}_2 \\ \text{HOCH}_2\text{CCH}_2\text{OH} \\ \text{NO} \\ \text{HOCH}_2\text{CHCH}_2\text{OH} \\ \text{LII} \\ \end{array} \begin{array}{c} \text{NO}_2 \\ \text{NO} \\ \text{NO} \\ \text{HOCH}_2\text{CCH}_2\text{OH} \\ \text{HOCH}_2\text{CH}_2\text{NO}_2 \\ \end{array} \begin{array}{c} \text{HOCH}_2\text{CNO}_2 \\ \text{NOH} \\ \text{LIII} \\ \end{array}$$

In accordance with the general reactivity of alkyl groups ortho and para to a nitro group, o- and p-nitrotoluene, nitro-p-xylene, o-nitroethylbenzene, m,p'-dinitrodiphenylmethane, and phenyl p-nitrobenzyl ether are nitrosated by amyl nitrite and an alkoxide. 138-141 Although there is not much published information about this reaction, it has been stated that the oxime of o-nitrobenzaldehyde (LIV) can be prepared with little difficulty if alcohol-free sodium ethoxide is used as catalyst.141

$$\begin{array}{c|c} CH_3 & CH=NOH \\ \hline NO_2 & C_5H_{11}ONO \\ \hline (N_BOC_2H_5) & \hline \\ LIV \end{array}$$

137 Earl, Ellsworth, Jones, and Kenner, J. Chem. Soc., 1928, 2697.

¹³⁹ Farbwerke vorm Meister, Lucius, and Brüning, Ger. pat. 107,095 [Chem. Zentr., 1900, I. 8861.

141 Lapworth, J. Chem. Soc., 79, 1274 (1901).

¹³³ Angeli and Angelico, Atti accad. nazl. Lincei, [5] 8, II, 28 (1899) [Chem. Zentr., 1899, II, 371].

¹⁴⁰ Farbwerke vorm Meister, Lucius, and Brüning, Ger. pat. 109,663 [Chem. Zentr., 1900, II, 458].

However, there is disagreement about the necessity of using alcohol-free sodium ethoxide in this reaction. 139

The activating effect of the nitro group described in the preceding paragraphs is to be contrasted with the opposite effect which this group sometimes exerts. Thus, although ethyl phenylacetate has been nitrosated successfully, 116 methyl 2,4-dinitrophenylacetate is reported to undergo nitrosation only in an alkaline medium. 16,121 p-Nitrobenzylmalonic acid and ethyl p-nitrobenzylacetoacetate are reported not to undergo nitrosation. 142

#### Hydrocarbons

As would be expected from the general reactivity of its methylene group, cyclopentadiene (LV) can be nitrosated in 70-90% yield. 143

HC=CH
$$\downarrow CH_2 \xrightarrow{C_2H_5ONO} HC=CH$$
HC=CH
$$\downarrow LV \qquad HC=CH$$
(as a dimer)

Lynn and his co-workers 144 found that sunlight catalyzes a reaction between hydrocarbons and nitrosyl chloride. Heptane was converted into the oxime of di-n-propyl ketone, 145 and toluene gave benzaldoxime in almost quantitative yield based on the nitrosyl chloride. 146

#### SYNTHETIC APPLICATIONS

#### α-Oximino Acids and Esters

 $\alpha$ -Oximino acids and esters are most frequently prepared by the nitrosation of substituted  $\beta$ -keto esters, malonic acids, and malonic esters. The other methods available for the preparation of these oximes are (1) reaction of an  $\alpha$ -keto acid or ester with hydroxylamine, 142, 147, 148, 149 (2) reaction of an  $\alpha$ -halo acid with hydroxylamine, 150 (3) reaction of an  $\alpha$ -halo ester with sodium nitrite, 92, 151, 152, 153 and (4) formation, oxidation,

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    Mattocks and Hartung, J. Am. Pharm. Assoc., 35, 18 (1946).
    Thiele, Ber., 33, 669 (1900).
    Lynn, J. Am. Chem. Soc., 41, 368 (1919).
    Lynn and Hilton, J. Am. Chem. Soc., 44, 645 (1922).
    Lynn and Addition. J. Am. Chem. Soc., 44, 645 (1922).
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in Lynn and Arkley, J. Am. Chem. Soc., 45, 1045 (1923). in Meyer and Janny, Ber., 15, 1525 (1882).

Puitti, Gazz. chim. ital., 17, 519 (1887).
 Erlenmeyer, Ann., 271, 167 (1892).

¹¹⁵ Hantzach and Wild, Ann., 289, 285 (1896).

Lepercq, Bull. soc. chim. France, [3] 9, 630 (1893).
 Lepercq, Bull. soc. chim. France, [3] 11, 295 (1894).

¹¹¹ Lepercq, Bull, soc. chim. France, [3] 11, 886 (1894).

and hydrolysis of an  $\alpha$ -hydroxylaminonitrile. The usefulness of reaction 1 is limited by the comparative unavailability of  $\alpha$ -keto acids, whereas reactions 2, 3, and 4 require relatively long reaction times.

α-Oximino acids and esters prepared by nitrosation reactions have been used extensively in the synthesis of the corresponding  $\alpha$ -amino acids and esters. The  $\alpha$ -amino acids which have been prepared in this manner are alanine, 92 α-amino-n-butyric acid, 92 α-amino-δ-diethylaminovaleric acid (ethyl ester), 8, 155 3,4-dihydroxyphenylalanine, 101 glutamic acid, 92, 93 β-hydroxyglutamic acid, 96 isoleucine, 92, 166 leucine, 12, 92 lysine, 157, 158 p-methoxyphenylalanine, 92 norleucine 92 (ethyl ester 8), norvaline, 92 phenylalanine 12, 92 (ethyl ester 8), the α-amino-β-hydroxyn-butyric acids, 159 and tyrosine. 92 In recent years many  $\alpha$ -amino acids have been prepared from substituted aminocyanoacetic and aminomalonic esters obtained from ethyl oximinocyanoacetate and diethyl oximinomalonate, respectively. 126, 160

 $\alpha$ -Oximino esters also provide a route to  $\alpha$ -keto acids and esters, since the oximino group can be replaced by a keto group by treatment with a nitrous acid derivative. 161-164 Diethyl oxomalonate (LVI) is prepared from diethyl malonate in 74 to 76% yield without isolation of the

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154 Miller and Plöchl, Ber., 26, 1545 (1893).
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Breslow, Walker, 1050, Bull. soc. chim. France, [3] 35, 965 (1906).

Bouveault and Locquin, Batter of the Bouveault and Locquin and Locquin and Locquin and Locquin and Locquin and Local Olynyk, Camp, Grinten, 13, 408 (1948).

188 Borsook, Deasy, Haagen-Smit, Keighley, and Lowy, J. Biol. Chem., 176, 1384 (1948).

¹⁵⁹ Adkins and Reeve, J. Am. Chem. Soc., 60, 1328 (1938).

¹⁶⁰ Albertson, J. Am. Chem. Soc., 68, 450 (1946).

¹⁵⁰ Albertson, J. Am. Chem. Soc. chim. France, [3] 31, 1142 (1904).
¹⁵¹ Bouveault and Locquin, Bull. soc. chim. France, [3] 31, 1142 (1904).

¹⁶² Kondo, Biochem. Z., 38, 408 (1912).

¹⁸³ Locquin, Bull. soc. chim. France, [3] 31, 1147 (1904).

¹⁶⁴ Sen, Biochem. Z., 143, 197 (1923).

$$CH_2(CO_2C_2H_5)_2 \xrightarrow{N_2O_3} CO(CO_2C_2H_5)_2$$
LVI

intermediate oximino ester. 165 The same reaction can be accomplished more satisfactorily by means of the commercially available nitrogen dioxide.166

The use of  $\alpha$ -benzyloximino acid chlorides in the synthesis of peptides is described on p. 271.

#### a-Oximino Ketones

α-Oximino ketones, prepared readily by the nitrosation of ketones and  $\beta$ -keto acids, have served in the synthesis of a large number of  $\alpha$ -diketones,  $\alpha$ -dioximes,  $\alpha$ -diamines,  $\alpha$ -amino alcohols,  $\alpha$ -amino ketones, and heterocyclic compounds. The diketones have been prepared in high yield by treatment of the  $\alpha$ -oximino ketones with dilute mineral acid 167 or with a nitrous acid derivative. 50, 168, 169 There is a report of the direct conversion, in high yield, of a \beta-diketone (p-nitrodibenzoylmethane) into the corresponding triketone by means of nitrous fumes. 50 Knorr's method 170,171,172 for the synthesis of pyrroles involves the reduction of an  $\alpha$ -oximino ketone to an  $\alpha$ -amino ketone, which, usually without isolation, is condensed with a ketone to form a substituted pyrrole. Ethyl acetoacetate is converted into 2,4-dimethyl-3,5-dicarbethoxypyrrole (LVII) by this procedure.¹⁷³ The amino ketones derived from α-oximino ketones

in Rielmomer and Irvine, Org. Syntheses, 25, 34 (1945).

in Kolb, Ann., 291, 280 (1896).

in Bouverult and Locquin, Bull. 20c. chim. France, [3] 31, 1169 (1904).

in Locquin, Bull ere, chim. France, [3] 31, 1173 (1904).

[&]quot;1 Knorr, Ann., 236, 317 (1886).

¹⁵ Ochisi, Tsuda, and Ikuma, Ber., 68, 1551 (1935).

er Ochia, Terda, and Ikuma, Ber., 68, 1710 (1935).

¹⁷ Fireher, Org. Syntheses, Coll. Vol. 2, 202 (1943).

have served also in the synthesis of imidazolones (LVIII) and thiolimidazoles (LIX). 33, 73, 174-177 The catalytic reduction of  $\alpha$ -oximino ketones

leads to  $\alpha$ -amino ketones,  $\alpha$ -amino alcohols,  $\alpha$ -hydroxy oximes, or pyrazines, depending upon the experimental conditions. 93,178 For example, the hydrogenation of ethyl α-oximinoacetoacetate (LX) over Raney nickel at 120 atm. yields, after oxidation of the product by air, 2,5-dimethyl-3,6-dicarbethoxypyrazine (LXI); hydrogenation at 320 atm.

furnishes ethyl  $\alpha$ -amino- $\beta$ -hydroxy-n-butyrate (LXII). Many of the  $\alpha$ -oximino ketones obtained from aryl alkyl ketones have been reduced to amino alcohols that have pressor activity. 21, 22, 31, 35, 37, 179, 180, 181

### EXPERIMENTAL CONDITIONS AND PROCEDURES

#### **Experimental Conditions**

Since nitrous acid derivatives can convert oximes into ketones, the nitrosation of aliphatic carbon atoms is usually carried out with only a small excess of nitrosating agent * and at temperatures between 0 and

- ¹⁷⁴ Fox, Sargent, and Buchman, J. Am. Chem. Soc., 67, 496 (1945).
- 176 Jackman, Klenk, Fishburn, Tullar, and Archer, J. Am. Chem. Soc., 70, 2884 (1948).
- 176 Ochiai and Ikuma, Ber., 69, 1147 (1936).
- 177 Wynn and Corwin, J. Org. Chem., 15, 203 (1950).
- Adkins and Shriner, in Gilman, Organic Chemistry, Vol. I, 2nd ed., p. 807, John Wiley
- 179 Glynn and Linnell, Quart. J. Pharm. Pharmacol., 5, 491 (1932). & Sons, New York, 1943.
  - 180 Hartung, Munch, and Crossley, J. Am. Chem. Soc., 57, 1091 (1935).
- 181 Machlis and Blanchard, J. Am. Chem. Soc., 57, 176 (1935). * An interesting exception to this practice is the conversion of methylhydrastein into its oximino derivative in 80% yield by means of a twenty-two fold excess of ethyl nitrite (see ref. 239).

50°. It is customary to add one reactant in small portions to a stirred solution of the remaining reactants.

The isolation of products is largely dependent upon the solvent and reagents employed. Oximes are frequently purified by extraction into sodium carbonate or sodium hydroxide solution, provided they are stable under these conditions. Since heating of oximino derivatives may cause violent decomposition, care should be used in attempts to distill these compounds or to remove solvents by distillation (see page 354).

It is not always possible to make a rigorous differentiation among the various reagents employed in the nitrosation of aliphatic compounds because the effective nitrosating agent is often formed after the reactants have been brought together. For example, ethyl nitrite is the nitrosating agent when it is used with sodium ethoxide as catalyst, but with hydrogen chloride as catalyst the agent is believed to be nitrosyl chloride ("nascent nitrosyl chloride").182 The following discussion is therefore based upon the reagents employed rather than on compounds believed to be formed in the reaction mixture.

The alkyl nitrites cause a marked fall in blood pressure by dilating the peripheral arteries. In large amounts they produce methemoglobinemia, resulting in cyanosis and asphyxia. Therefore, alkyl nitrites, particularly methyl and ethyl nitrites, which are gases at room temperature, should be used with caution

- 1. Inorganic nitrite and acid. This combination possesses the advantage of avoiding the preliminary preparation of the nitrosating agent. It can be used with both water-soluble and water-insoluble compounds. The water-insoluble compounds have been nitrosated by employing glacial acetic acid as solvent and sodium nitrite dissolved in the minimum amount of water. Nitroparaffins are usually nitrosated by the addition of nitrite and mineral acid to an alkaline solution of the nitro compound.
- 2. Alkyl nitrite and an alkoxide. This effective combination is almost always used in ethanol solution. Only in the conversion of o- and pnitrotoluene to the corresponding benzaldehyde oxime has alcohol-free alkoxide been said to be necessary,141 but even in this case there is a conflicting report.119 The claim 29 that the presence of a trace of water increases the yield of diacetylmonoxime from methyl ethyl ketone could not be confirmed 12

Ethyl nitrite nitrosates camphor more readily and in higher yield than does amyl nitrite.14 Sidgwick 14 attributes to Slater 22 a report of

to Rivenboldt and Schmitz-Dumont, Ann., 444, 113 (1925).

¹¹ Semen and Damerell, J. Am. Chem. Soc., 47, 2033 (1925). in Rupe and Spintrenber, Ber., 40, 4313, footnote (1907).

the more rapid action of methyl and ethyl nitrites as compared to amyl nitrite; and, although no statement or experiment regarding this question appears in the paper by Slater, it is probably true that the lower homologs are more reactive. Probably a more important advantage of the use of one of the gaseous nitrites is that the alcohol (methanol or ethanol) formed in the reaction is both miscible with water and readily volatile, so that its presence does not complicate the isolation of the product. Gero and Seitchik ^{184a} recommend n-propyl nitrite as also having this advantage and as being preferred to methyl and ethyl nitrites because it can be handled as a liquid (b.p. 46-49°).

- 3. Alkyl nitrite and hydrogen chloride. This is the most widely used reagent combination. It has the advantage of yielding a reaction mixture which, by vacuum distillation, can be freed of reagents and at least one by-product, the alcohol formed from the nitrite. Ethanol and ether are used most frequently as solvents. A small amount of concentrated hydrochloric acid is often the source of the hydrogen chloride, but many nitrosations are carried out under anhydrous conditions. Slater 32 and Aston and Mayberry 19 found that water decreased the activity of the catalyst in ketone nitrosations, but Semon and Damerell 183 reported that a small amount of water had very little effect on the yield of diacetylmonoxime from methyl ethyl ketone. In a study of the nitrosation of a number of phenacyl chlorides, it was found necessary to add a trace of water to initiate the reaction of p-methoxyphenacyl chloride with isopropyl nitrite and hydrogen chloride.39 With some ketones, maximum yields of their oximino derivatives depend upon the use of an optimum concentration of catalyst. 32, 183 Many nitrosations require continuous introduction of hydrogen chloride, but a trace suffices in the reaction between ethyl nitrite and α-acetyl-γ-butyrolactone.95 though the use of a large amount of hydrochloric acid has been reported to lead to nitrosochlorination, 27,185 normally the use of an alkyl nitrite and hydrogen chloride is not complicated by this side reaction. Acetyl chloride can be used in place of hydrogen chloride.19
- 4. Nitrosation in concentrated sulfuricacid. Bouveault's method  $^{10,88-91}$  employing nitrosylsulfuric acid ("lead chamber crystals") in concentrated sulfuric acid for the nitrosation of  $\alpha$ -substituted  $\beta$ -keto esters has been replaced by the more convenient method of Hartung,  9,92  in which nitrosation is accomplished by n-butyl nitrite in 85% sulfuric acid. Its usefulness depends upon the stability of the compounds employed in the strong acid.  9

¹⁸⁴a Gero and Seitchik, private communication.

¹⁸⁵ Claisen and Manasse, Ann., 274, 95 (1893).

- 5. Nitrosyl chloride. The use of this reagent is attended with the disadvantage that nitrosochlorination as well as simple nitrosation may occur. 182, 186, 187
- 6. Nitrous fumes. This reagent is seldom used at present for the nitrosation of aliphatic carbon atoms. It has had, however, extensive use in the preparation of N-nitroso-N-acetylarylamines.^{187a} The fact that the reagent is a gas as well as a mixture of nitrogen oxides makes it difficult to employ it in a quantitative manner.

#### Experimental Procedures

The preparations of methyl nitrite,³⁶ ethyl nitrite,¹⁵ and *n*-butyl nitrite ¹⁸⁸ are described in *Organic Syntheses*. *n*-Butyl nitrite and, presumably, other organic nitrites decompose after several weeks at room temperature.

Directions for the preparation of oximinoacetone, diacetyl monoxime, 2-oximino-3-pentanone, and 2-oximino-3-hexanone are given by Fischer and Orth.^{185a}

Detailed procedures for the preparation of diacetyl monoxime,  $\alpha$ -oximinopropiophenone, and phenylglyoxylohydroxamyl chloride ( $\omega$ -chloroisonitrosopropiophenone) from the corresponding ketones in yields of 69-74%, 65-68%, and 82-86%, respectively, are given in Organic Syntheses. 15, 36, 40 Alkyl nitrites and hydrogen chloride or hydrochloric acid are used to effect the nitrosations.

Dioximinoacetone from Acetonedicarboxylic Acid. A solution of 150 g. of crude acetonedicarboxylic acid  188b  in 275 ml. of water is cooled in an ice-salt bath. A solution of 100 g. of sodium nitrite in 200 ml. of water is added slowly, with stirring, while the temperature of the reaction mixture is kept below 0°. The mixture is cooled to  $-5^{\circ}$  and filtered immediately. The solid is washed with small portions of ice water. An additional amount is obtained by adding 200 ml. of cold 6 N nitric acid to the filtrate. The white product is washed with four small portions of ice water and dried over sulfuric acid in a vacuum desiccator. The product weighs 59 g. (51%) and decomposes at 133°.

¹⁸c Demole, Ann., 175, 146 (1875).

¹² Rheinboldt and Schmitz-Dumont, Ber., 61, 32 (1928).

[&]amp; Sons, 1944.

¹⁴³ Noyes, Org. Syntheses, Coll. Vol. 2, 108 (1943.)

¹⁹⁴ H. Fischer and H. Orth, Die Chemie des Pyrrole, Vol. 1, pp. 408-410, Akad. Verlag, Leipzig, 1934.

acetonedicarboxylic acid contains sulfuric acid.

3-Oximino-5-ethoxy-2-pentanone from Ethyl  $\alpha$ -2-Ethoxyethylaceto-acetate. To 20 g. of 5% sodium hydroxide solution is added 78 g. of ethyl  $\alpha$ -2-ethoxyethylacetoacetate, and the mixture is stirred for nine hours. Then 26.6 g. of solid sodium nitrite is added, and the orange solution is cooled in an ice bath while a solution of 30 ml. of concentrated sulfuric acid in 80 ml. of water is slowly added from a dropping funnel. The solution is allowed to stand overnight. It is then made alkaline with 10% sodium hydroxide solution and extracted with ether. The aqueous solution is acidified with sulfuric acid (saturation with carbon dioxide may be used), the product separating as a red-brown oil. The aqueous layer is extracted with ether, and the combined oil and extracts are washed free of acid, dried over sodium sulfate, and distilled. The yield of product boiling at 108–113°/2.3 mm. is 30 g. (49%). The freezing point of a redistilled sample (116–116.5°/1.4 mm.) is 29.5°.

Ethyl  $\alpha$ -Oximinoacetoacetate from Ethyl Acetoacetate. Is In a 5-1. three-necked flask fitted with a thermometer, a reflux condenser, and a mechanical stirrer are placed 730 ml. (750 g., 5.8 moles) of commercial ethyl acetoacetate and 840 ml. of glacial acetic acid. The flask is cooled in an ice-salt bath, and a solution of 450 g. of 95% sodium nitrite in a liter of water is added over a period of approximately one hour, the temperature being kept at 25°. Three liters of water is then added, and stirring is continued for two hours.

One quarter of the reaction mixture is placed in a 2-1. separatory funnel and shaken with 350 ml. of ether. The bottom aqueous layer is run off, and the next quarter of the reaction mixture is placed in the separatory funnel and extracted with the same ether. This is repeated until all is extracted. This cycle is repeated twice, using 200 ml. of ether each time. The ether extracts are combined, washed once with water, four times with sodium bicarbonate solution, and once more with water. The addition of sodium chloride is occasionally necessary to cause the layers to separate promptly. After drying the ether solution with sodium sulfate, the solvent is distilled on a steam bath at atmospheric pressure, and then for two hours at about 35 mm. The residue of brown, liquid, impure ethyl  $\alpha$ -oximinoacetoacetate weighs 650-700 g.

The crude product is dissolved in toluene (120 ml. per 100 g. of crude material) and the solution is filtered. Cooling to  $-13^{\circ}$  to  $-15^{\circ}$  with stirring for one-half hour causes crystallization. The solid is filtered, washed with a little cold toluene, and air dried overnight. A yield of 550-600 g. (63%), m.p.  $57.5-58^{\circ}$ , is obtained. The addition of petroleum

¹⁸⁹ Tota and Elderfield, J. Org. Chem., 7, 317 (1942).

ether (b.p. 60-90°) to the toluene decreases the solubility of the product and permits an increased yield (75%). 150

The once-crystallized material may be recrystallized from toluene, but 180 ml. of solvent should be used per 100 g. of oximino ester; the recovery of pure white product, m.p. 58-58.5°, is 90%. If the toluene mother liquor is distilled on a hot plate at atmospheric pressure, the oximino ester decomposes, sometimes violently. The mother liquor can be used for crystallizing the next batch of crude material, or most of the toluene can be distilled under reduced pressure on a steam bath and 50-60 g. more of the oximino ester, m.p. 56°, obtained on cooling.

α-Oximino-γ-butyrolactone from α-Acetyl-γ-butyrolactone. To a cold (0° to -5°) solution of 256 g. (2 moles) of α-acetyl-γ-butyrolactone in 500 ml. of methanol is added 300 g. (4 moles) of ethyl nitrite. The reaction flask is packed in ice and salt and allowed to stand for fifteen to twenty hours, during which time the ice melts and the temperature reaches that of the room. The mixture is cooled, and the crystalline solid is collected on a filter. The filtrate is concentrated under diminished pressure, and the dark-colored residue is heated on the steam bath with 100 ml. of n-butyl alcohol. The mixture is cooled and filtered. The two crops of crystals are combined, washed twice with 100-ml. portions of cold n-butyl alcohol and then with ether. The α-oximino-γ-butyrolactone weighs 196-209 g. (85-91%) and melts at 183-185° (lit. 192°).

α-Oximinocaproic Acid from Ethyl n-Butylacetoacetate.9 In a 400-ml. beaker surrounded by an ice-salt bath is placed 30 g. of 85% sulfuric acid. Mechanical stirring is started (a four-blade paddle stirrer was found most efficient), and, when the temperature of the acid reaches  $-5^{\circ}$  to  $0^{\circ}$ , 18.6 g. (0.1 mole) of ethyl *n*-butylacetoacetate is added slowly enough that no rise in temperature occurs. When this addition is complete, 11 g. (0.105 mole) of n-butyl nitrite is slowly added dropwise, with the temperature as near 0° as possible. Slow effervescence is observed, but, if the nitrite is added too rapidly, oxides of nitrogen are evolved. After all the nitrite has been added, small pieces of ice are added to dilute the acid. At this point a white, curdy precipitate of oximino ester appears. Cold water is then added, and the liquid is extracted with ether. The oximino compound is extracted from the ether by cold 10% sodium hydroxide solution. The red alkaline extract is heated on the steam bath for fifteen minutes, then cooled and acidified. The precipitated  $\alpha$ -oximinocaproic acid is filtered, and the filtrate is

¹⁹⁰ Albertson, Tullar, King, Fishburn, and Archer, J. Am. Chem. Soc., 70, 1150 (1948).
* Evidently the reaction is catalyzed by a trace of hydrogen chloride present in the ethyl nitrite, since with ethyl nitrite prepared from sulfuric acid the reaction proceeds very slowly unless a small amount of an acid is added.

extracted with ether. The product is recrystallized from petroleum ether and melts at 136° (dec.); the yield is 12.5 g. (86%).

By the same procedure, ethyl  $\alpha$ -benzylacetoacetate is converted into  $\alpha$ -oximino- $\beta$ -phenylpropionic acid in 85% yield. No oxime could be obtained from ethyl 3,4-diethoxybenzylacetoacetate by this procedure.

Ethyl α-Oximinocaproate from Diethyl n-Butylmalonate.8 Sixty-four and nine-tenths grams (0.3 mole) of diethyl n-butylmalonate is placed in a 500-ml. flask equipped with a mercury-sealed stirrer, dropping funnel, and an ice-water-cooled condenser carrying a drying tube. The flask is immersed in an ice bath, and 33.8 g. (0.4 mole) of ethyl nitrite * is added to the stirred solution, the temperature of which is maintained at about  $0^{\circ}$ . The mixture is then cooled to  $-10^{\circ}$  in an ice-salt bath, and a solution of sodium ethoxide (prepared from 6.9 g. of sodium and 138 ml. of absolute ethanol) is added slowly with stirring. The flask is stoppered tightly and kept in a freezing unit of a refrigerator at -10° for twelve hours. The mixture is poured into an evaporating dish which is kept in a vacuum desiccator over concentrated sulfuric acid until the alcohol has evaporated. (The alcohol may be removed rapidly with equally good results by gently heating the mixture on a steam bath under reduced pressure.) To the residue is added an equal volume of ice water, and the aqueous solution is extracted with ether.† While it is cooled in an ice bath, the aqueous solution is acidified to pH 5 with cold concentrated hydrochloric acid. (During the neutralization, ice is added directly to the aqueous solution.) The  $\alpha$ -oximino ester, which precipitates as a yellow oil, is taken up in ether, and the aqueous solution is extracted several times with ether. The combined ether extracts are dried over Drierite, and the solvent is distilled, leaving 42.8 g. (83%) of ethyl α-oximinocaproate as a light yellow solid, m.p. 49-53°. Recrystallization from petroleum ether (b.p. 30-60°) yields 41.4 g. (80%) of a white product melting at 53-55°.

By a similar procedure, diethyl benzylmalonate is converted into ethyl  $\alpha$ -oximino- $\beta$ -phenylpropionate in 92% yield.

^{*} Purified commercial butyl nitrite gave quite impure ethyl  $\alpha$ -oximinocaproate.

[†] From the ether solution, after drying and removing the solvent, there was obtained 9.4 g. (27%) of diethyl carbonate.

#### TABULAR SURVEY

The data in the tables cannot always be used to determine the superiority of a particular nitrosation procedure inasmuch as many preparations were not carried out with a view to obtaining maximum yields. Experimental procedures have been indicated by the following notations.

 $HNO_2$  = sodium nitrite and mineral or acetic acid.

N₂O₃ = nitrous fumes evolved from a mixture of concentrated nitric acid and arsenic trioxide.

NOCl = nitrosyl chloride.

NO₂SO₃H = nitrosylsulfuric acid in concentrated sulfuric acid.

 $C_4H_9ONO_1$  85%  $H_2SO_4 = n$ -butyl nitrite in 85% sulfuric acid.

RONO, HCl = alkyl nitrite * and hydrogen chloride. (Differentiation between anhydrous hydrogen chloride and concentrated aqueous hydrogen chloride has not been made unless the two reagents were compared under similar conditions.)

RONO, CH₃COCl = alkyl nitrite * and acetyl chloride.

RONO, MOR = alkyl nitrite * and an alkoxide.

HOH = hydrolysis.

Where more than one reference is given for a single entry, the yield reported is taken from the reference in italics.

Although many examples of the reaction are not listed in abstract journals, it is hoped that practically all those recorded in the literature prior to the January, 1950, issue of *Chemical Abstracts* † have been detected. A number of more recent examples are also included in the tables. The compounds are in general listed in order of increasing size and complexity, particularly as regards the group which is nitrosated. Methyl ketones therefore precede other dialkyl ketones, which are in turn followed by alicyclic ketones and then aryl alkyl ketones. In each of the tables, examples of the nitrosation of methyl groups precede examples of the nitrosation of methyl groups.

^{*}Amyl nitrite and isoamyl nitrite are both listed as C5H11ONO because commercial products may be mixtures of isomers.

[†] For the convenience of the reader, Chemical Abstracts references have been included for several foreign articles listed in this chapter. However, except for references 26, 235, 237, and 248, the original papers have been consulted.

## TABLE

	% Keieronos	23 191	_	91 69	40 32		1		191	10	97	63, 69-74 183, 15			62 198, 199		58		31 + 183, 187, 193		202	30-70		8	£ .	2. C.				_	Ž.	61 ·	
Kerones	Products	A. Dialkyl Monoketones	Acetylmethyl nitrolic acid *	Janitrosodincetone nitrate(?)	Oximinoacetone	Oximinoncetone	Oximinoacctone	Oxininoacetone, chlorocximinoacetone, nitroacettonia	tion products of phorone	Diacetyl monoximo	Ethyl nitrolie acid	Diacetyl monoxime	Diacetyl monoxime	Dincetyl monoxime		Diacetyl monoxime	Diacetyl monozime	Directyl monoxime	Diacetyl monoxime	Diacetyl monoxime	Diacetyl monoxime	Discetyl monorime	3-Oximino-2-pentanone il	3-Nitroso-3-methyl-2-butanone	3-Nitroso-3-methyl-2-butanone	3-Nitroso-3-methyl-2-butanone	3-Orimino-5-pentanol-2-one	3-Oximino-2-hexanone	Methyl S-oximino-1-keto-2-pentenonte	1-Oximino-f-methyl-3-penten-2-one	2.Oximino-2-octanone	Mathyl anitrosocycloheryl ketone	
	Method	A.		$N_2O_3$	N ₂ O ₃	HNO ₂	CHJONO, HCI	C'HIONO, HOL	NOC	9	10.03	CH-ONO HC	O'C'H'C	CHILONO HO	Carrie of the ca	C.H.,ONO. HCI	C.H.,ONO. NaOCaHs	C.H.,ONO. NaOH	CAHIONO	CON	H-OS-ON	# HOSON	C.H. ONO. HCI	CHLONO HCI	CONO. D. HCI	CTUP CHOOL	CZHSONO, CHICAGO	Cthjono, iici	Cannono, noi	Cansolvo, nel	Canilono, nacosara	Csnilono, noi	
		Starting Compound			Acelone						Methyl ethyl ketone													Methyl n-propyl ketone	Methyl isopropyl ketone			Acetopropyl alcohol	Methyl $n$ -hutyl ketone	Methyl 4-keto-2-pentenoate	Mesityl oxide	Methyl n-bexyl ketone	

			67	ç
Methyl cyclohexyl ketone $(Cont^{\prime}d)$	C2H SONO, CH SCOCI	Methyl c-nitrosocyclonexyl ketone	2	CT :
Methyl benzyl ketone	NOCI	1-Oximino-1-phenyl-2-propanone	l	182
	CsH110NO, NaOC2Hs	1-Oximino-1-phenyl-2-propanone	92	167
4-Phenvl-2-butanone	C, H110NO, NaOC2H5	3-Oximino-4-phenyl-2-butanone	1	202
Benzalacetone	CsH110NO, HC1	1-Oximino-4-phenyl-3-huten-2-one	30-70	34,205a
Anisalacetone	NOCI	1-Oximino-4-p-methoxyphenyl-3-huten-2-one	ì	182
Methyl n-nonyl ketone	CsH110NO, HC1	3-0ximino-2-undecanone	30	206
2,4-Dinitrophenylacetone	CsH110NO, HC1	1-0ximino-1-(2,4-dinitrophenyl)-2-propanone	80	16
	CsH110NO, NaOC2Hs	3-Acetyl-6-nitrobenzisoxazole	ı	16
Diethyl ketonc	Control HCI	2-Oximino-3-pentanone	37-55	207
	C ₆ H ₁₁ ONO, HCI	2-0ximino-3-pentanone	30-70	34
Ethyl n-propyl ketone	C ₅ H ₁₁ ONO, HCl	2-Oximino-3-hexanone, 4-oximino-3-hexanone	ì	17
Ethyl isopropyl ketone	C2H5ONO, HC1	2-Oximino-4-methyl-3-pentanone	53	19
		2-Nitroso-2-methyl-3-pentanone	30	
	C ₂ H ₅ ONO, aq. HCl	2-Oximino-4-methyl-3-pentanone	27	20
		2-Nitroso-2-methyl-3-pentanone	7	
	C2H3ONO, CH3COCI	2-Oximino-4-methyl-3-pentanone	34	19
		2-Nitroso-2-methyl-3-pentanone	49	
	C ₅ H ₁₁ ONO, NaOC ₂ H ₅	2-Oximino-4-methyl-3-hutanone	40	208
Ethyl n-butyl ketone	C,H110NO, HCI	2-Oximino-3-heptanone, 4-oximino-3-heptanone	١	17
Ethyl isobutyl ketone	C ₆ H ₁₁ ONO, NaOC ₂ H ₅	2-Oximino-5-methyl-3-hexanone	40	208
Ethyl n-amyl ketone	C ₆ H ₁₁ ONO, HCl	2-Oximino-3-octanone, 4-oximino-3-octanone	i	17
Ethyl 180amyl ketone	C ₅ H ₁₁ ONO, HCl	2-Oximino-6-methyl-3-heptanone	١	17
Ethyl isonexyl ketone	C ₅ H ₁₁ ONO, HCl	2-Oximino-7-methyl-3-octanone	١	17
Ethyl cyclonexyl ketone	C2H,ONO, HC1	2-Oximino-1-cyclohexyl-1-propanone	33	19
		1-(1-Nitrosocyclohexyl)-1-propanone	4	
	C2H5ONO, aq. HCl	2-Oximino-1-cyclohexyl-1-propanone	15	19
		1-(1-Nitroscoyclohexyl)-1-propanone	4	
	C2H50NO, CH3COCI	2-Oximino-1-cyclohexyl-1-propanone	26	19
Ethyl n-pentadecyl ketone	OH ONO H	1-(1-Nitrosocyclohexyl)-1-propanone	က	
	Osmilono, noi	2-Oximino-3-octadecanone	1	18
Mater Deference 101 010				

Note: References 191-316 are listed on pp. 375-377.

* This compound was obtained as an oil of 52% purity. The yield was hased upon the weight and analysis of the oil. † The yield was based on the dioxime isolated.

‡ This yield could not he obtained in a confirmatory study; see ref. 183.

§ The solvent was concentrated bydrochlorie acid rather than sulfuric acid.

This was originally believed to be 1-oximino-2-pentanone. Kalischer, Ber., 28, 1513 (1895) This compound decomposes slowly when stored in air.

# TABLE 1—Continued

## Ketones

	Reference	82	209	193	19	19	19	10	10		19		ů.	•	ě	<b>7</b> 6	100	210	61.	211	211	212		63	211	61	63	62	62	61	184	213, 214
Yield	%	1	1	i	47	25.	66	3 5	6	25			, 0	•	200	Small	ļ	}	1 8	69	1	88		1	1	l	-	8	20	Small	(48)	32, 34
Ketones	Products	A. Dialkyl Monokelones-Continued	2-Oximino-3-eicosanone	3-Oximino-4-heptanone	3-Oximino-4-heptanone	4-Oximino-2-niethyl-3-hexanone	4-Oxinnino-2-methyl-3-hexnnone	4-Oximino-2-methyl-3-heptanone	4-Oximino-2-methyl-3-heptnhone	4-Oximino-2,5-dimethyl-3-hexanone	2-Nitroso-2,5-dimethyl-3-neximone	4-Oximino-2,5-dimethyl-3-hexhnone	2-Nitroso-2,5-dimethyl-3-hexanone	4-Oximino-2,5-dimethyl-3-hexanone	2-Nitroso-2,5-dimethyl-3-hexnnone	1,3-Dioximino-1,3-diphenyl-2-propanone	2,5-Dioximinoeyelopentanone	I,2-Cyclohexanediono monoximo nitrite(?)	2,6-Dioxininoeyelohexanone	1,2-Cyclohexanedione monoximo				3-Methyl-2,6-dioximinocyelohexanone	2-Oximino-4-nethyleyelohexnnone	2-Oximino-4-methyleyelohexnnone	2,6-Dioximino-3-methyleyelohexanone	3,5-Dioximino-2,2,6-trimethyl-4-piperidone	3,5-Dioximino-2,2,6,6-tetramethyl-4-piperidono	2.7-Dioximinoeyeloheptanone	o-Oximinocumphor	a-Oximinocam phor
	Method	A. Dialkyl .	CAHAONO, HCI	C.H.,ONO, IICI	NOON	C.H.ONO, HCI	Callagno, CHacoci	C.HSONO, HCI	C2H3ONO, CH3COCI	C2H6ONO, IICI		C ₂ II ₅ ONO, nq. HCl		Calfagno, Ciliacoci		C2H3ONO, NnOC2H5	CsH,10NO, CH,COCI	Callaono, HCI	Chilono, CH ₃ COCI	(+,-)-2-Octyl nitrite,	NaOC2115	(+)-2-Octyl nitrite, NaOC2tas	NaOC.Hs	CsHuono, CHacoci	(+)-2-Octyl nitrite, NaOC2Hs	(-)-2-Octyl nitrite, NaOC2Hs	CsH110NO, CH1COCI	C,H,,ONO, HC	C,H,,ONO, IICI	C,H,,ONO, CH,COCI	Chichen Chiche	Chilono, NaOC2Hs
	Starting Compound		American Land Land	Ethyl n-hephadeeyi ketano	Disn-propyi ketene	and the second lives of	n-I'ropyi isahiapyi vetoiic	Toursell ashutul ketene	success of the state of the sta	Jeonropyl isobutyl ketone						Dibenzyl ketone	Cyclopentanene	Cyclebexprope	Cyclonexamone					3-Methylevelehexanone	4-Methylevelohexanone		6-Oximino-3-methylevelohexanone	2.2.6-Trimethyl-4-piperidene	2.2.6.6-Tetramethyl-4-piperidone	Cyclobentanene	Compher	

6-Phenyleamphor	C5H110NO, NaNH2, KNH2	3-Oximino-6-phenyleamphor	1	215
(-)-Epieamphor	C ₆ H ₁₁ ONO, NaNH ₂	(-)-3-Oximinoepicamphor	71 (Cr.)	09
(+)-Carone	C5H11ONO, CH3COCI	(+)-Nitrosocarone	45	216
Menthone	C,H110NO, HC1	β,}-Dimethyl-←oximinocaprylio acid	Poor	59
	Chilono, HCI	4-Nitrosomenthone	œ	58
		β, ?-Dimethyl-ε-oximinocaprylio acid	99	
	C2H5ONO, CH3COCI	4-Nitrosomenthone	40	57
	C5H11ONO, NaOC2H6	8,2-Dimethyl-e-oximinocaprylic acid	89	59
Pulegone	Cohilono, HCI	2-Nitrosopulegone	1 10	217, 218
	Chilono, NaOC2Hs	β, t-Dimethyl-e-oximino-7-oetenoio acid	í	59
Dihydrocarvone hydrobromide	C2H6ONO, CH3COCI	1-Nitrosodihydrocarvone hydrobromide	15	57
Dihydroeucarvone	C ₅ H ₁₁ ONO, HCl	Nitrosodihydroeuenryone	1	216
	HNO ₂	Nitrosodihydroeucarvone	1	219
Carvomenthone (tetrahydrocarvone)	C2HbONO, CH3COCI	1-Nitrosocarvomenthone	-, 18	57, 220
Lropinone	Chilono, HCl	a,a'-Dioximinotropinone	06	65
	CaHuono, NaOCaHs	a,a'-Dioximinotropinone	ì	65
cts-tn-Acetyl-7-keto-8-methyldecahydroiso- quinolino	C2H4ONO, NaOC2H6	N-Acetyl-10-oximinodihydrohomomeroquinene ethyl	89	9
		ester		
		B. Aryl Alkyl Monoketones		
Aeetophenone	CH30NO, HCI	a-Oximinoacetophenone	** 69	32
	C,H,ONO, HCI	a-Oximinoacetophenone	6-12	31
	C4H3ONO, NaOC2H5	a-Oximinoacetophenone	-, 37	31, 21
	C ₅ H ₁₁ ONO, N ₁₁ OC ₂ H ₅	a-Oximinoacetophenone	06	28.30
Tables of the state of the stat	CoH110NO, NaNH2	a-Oximinoacetophenone	32	23
T-Brown october 1	C4H9ONO, HCI	a-Oximino-m-bromoacetophenone	75	32
m-Chlorogetonhonon	C4H9ONO, NaOC2H5	a-Oximino-p-bromoacetophenone	63	58
7-Chlorosestonbenone	C4H3ONO, HCI	α-Oximino-m-chloroacetophenone	63	32
amorrandonamorran	C4H90NO, NaOC2Hs	a-Oximino-p-chloroaectophenone	09	35
	CsH110NO, NaOC2Hs	α-Oximino-p-chloroacetophenone	l	221
3.4-Dichloroacetonhenone	NOC!	a-Oximino-p-chloroacetophenone	1	182
	Cannono, Na	a-Oximino-3,4-dichloroacetophenone	ī	179
3-Chloro-4-hydroxyacetonhenone	CHIONO, NAOC2HS	a-Oximino-3,4-dichloroacetophenone	51	179
	CAMBONO, MACCAES	a-Oximino-3-chloro-4-hydroxyacetophenone	ৠ	21
3-Bromo-4-hydroxyacetophenone	CARSONO, INC.	a-Oximino-3-chloro-4-hydroxyacetophenone	11	21
	Offigoro, no.	a-Oximino-3-bromo-4-hydroxyaeetophenone	i	21
Note: References 191-316 are listed on pp. 375-377.	375-377.			

Note: References 191-316 are listed on pp. 375-377.
** The yield was calculated on the basis of unrecovered ketone.

# TABLE I-Continued

	Reference	222	23	223	224	224	225	225	39, 40	08	8 8	£0	30	39	39	39	16	226	36, 32	35, 37	33.87	200	37 180	201.00	000	006	077	827	5.7	24	24	24	24	24, 180	
Yield	%	ļ	75	7.1	1	l	44	: 1	98 60	06, 00	# 6	82	7.2	87	95	85	52	! !	65_68 75	51 72	30-70	** ** **	74 70	0,4,	64, 88	2 }	2	75	74	87	88	92	83	83, 89	
Ketones	Products	B. Aryl Alkyl Monoketones-Continued	a-Oximioo-p-methylacetophenone	a-Oximino-3,4-dimethoxy acceptance	a-Oximino-p-benzyloxyacctophenone ++	a-Oximino-2-benzyloxy-o-methoxymetalynemin	a-Oximino-2-benzyloxy-5-ethoxynectophenone 11	a-Oximinophenacylpyridinium bromide	a-Oximinophenacylpyridinium bromide	Phenylglyoxylohydroxamyl chloride	n-Tolvielvoxvlobydroxamyl chloride	Vanilalinarilohydroxamyl chloride	Ottombonical characteristics of the character	poniorophical designation of the contract of t	p-Methoxyphenyiglyoxylong arcamon photoide	p-Hydroxyphenyigiyoxyjoniyaroxamiy oning	3,4-Dihydroxyphenylglyoxylobydroxamyl culonide	Phenyl a-oximino-2,4-dinitropenzyl ketone	a-Oximino-a-2-quinolyi-o-carboxyacecophenome	a-Oximinopropiophenone	a-Oximinopropiophenone	a-Oximinopropiophenone	a-Oximinopropiophenone	a-Oximino-p-methylpropiophenone	a-Oximino-p-phenylpropiophenone	a-Oximino-p-nitropropiophenone	a-Oximino-p-acetamidopropiophenooe	a-Oximino-p-benzamidopropiophenone	a-Oximino-o-fluoropropiophenone	a-Oximino-m-fluoropropiophenone	Oriming m-fluoropropionismone	-Orimino-ephononionhenone	- Orimino - hloroprophonone		a-Oximino-p-eniorapiopiopiopio
	Method	B. Aryl.	C ₅ H ₁₁ ONO, NnOC ₂ H ₆	C,H110NO, NaOC2H5	C,H,ONO, NAOC,H,	CH.ONO IICI	CH,ONO, HC	Cramono	HNO	TONO HCI	ONO II	CHOON, HO	C,H,ONO, HC	C,H,ONO, HC	C4H,0NO, HCI, H20	C,H,ONO, HC!	C,H,ONO, HCI	C,H,10NO, HC	HNO2	CH ₂ ONO, HCl	C,H,ONO, HCl	C,H110NO, HCI	C,H110NO, NaOC2H5	C'HONO, HCI	C,H,ONO, HCI	C,H,ONO, HCI	C, HONO, HCJ	C'H'ONO HO	CH CNCHO	CHOOLECT	ONO HO	Citationo, inci	CARGONO, HOL	CHRONO, HOL	C,HOONO, HCI
	Starting Compound		and market and an artist and	p-Metayincetopmenture	3,4-Dimethoxyntetophenone	p-Henzyloxyacetophenoue	2-Benzyloxy-5-methoxyacetophenone	2-Renzyloxy-5-ethoxyacctophenoue	Phenacylpyridinium bromine		Phenacyl chloride	p-Methylphenacyl chloride	p-Phenylphennoyl chloride	r-Chlorophenaeyl chloride	n-Methoxyphenacyl chloride	Tridroganhengen chloride	3 4-Dihydroxynhenseyl chloride	Phenyl 2.4-dinitrobenzyl ketone		Pronionhenone				m. Mother forcation honone	The property of the property o	- Nitrontonionlenone	A actomidononionhonono	p-Accumidontopiopione	P.Denkinnacjivopiopiono	ortinosopiopiopionies	m-rinordordordordordordordordordordordordordo	p-r luoroprophenone	e-Chloropropiophenone	m-Chloropropiophenone	p-Chloropropiophenone

NIIROSATION OF ME	
	<i>\$</i>
35 24 24 35 35 35 22 22 22 22 22 22 22 22 22 22 22 22 22	231 31 31 31 31 31 31 31 31 232 232 234, 236 180 180 236
Good 71 76 82, 87 1 \$\$ 41 72 90 50-75 50-75 50-75 50-75 72 88 60	50 50 81 81 69 55-60 25 45 14 74
α-Oximino-p-chloroptopiophenone α-Oximino-p-chloroptopiophenone α-Oximino-p-tromopropiophenone α-Oximino-p-tromopropiophenone α-Oximino-p-tromopropiophenone α-Oximino-p-methoxypropiophenone α-Oximino-2,-d-dinethoxypropiophenone α-Oximino-2,-d-dinethoxypropiophenone α-Oximino-2,-d-dinethoxypropiophenone α-Oximino-2,-b-arayloxypropiophenone α-Oximino-2,-b-arayloxypropiophenone α-Oximino-2,-b-arayloxypropiophenone α-Oximino-2-b-arayloxypropiophenone α-Oximino-2-b-arayloxypropiophenone α-Oximino-2-c-thylearhonsto-5-methoxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone α-Oximino-3-hydroxypropiophenone	Acetophenone oxime  α-Oximino-β-phenylpropiophenone α-Oximino-p-methylhutyrophenone α-Oximino-p-methylhutyrophenone α-Oximino-o-carbomethoxyhutyrophenone α-Oximinovalerophenone α-Oximinovaprophenone α-Oximinocaprophenone α-Oximino-1-acetonaphthone α-Oximino-1-nectonaphthone α-Oximino-1-propionaphthone α-Oximino-1-propionaphthone β-oximino-2-propionaphthone β-oximino-2-propionaphthone
C,H,ONO, HCI C,H,ONO, HCI	CAHAONO, NAOCARS CAHIONO, NAOCARS CAHIONO, NAOCARS CAHONO, HCI
p-Chloropropiophenone (Gont'd) o-Bromopropiophenone m-Bromopropiophenono p-Bromopropiophenono g-Jilano-1-hydroxypropiophenone g-Ji-Dimethoxypropiophenone	3,4-Dihydroxypropiophenone  a-Pitenylpropiophenone  GH10N  β-Pitenylpropiophenone  GH50N  Huyrophenone  C-Arbomethoxybutyrophenone  C-Arbometh

[#] The oxime was reduced, without isolation, to the dehenzylated amino ketone-Note References 191-316 are listed on pp. 375-377.

^{\$\$} The yield was based on the amino ketone isolated. :: The yield was based on the dioxime isolated.

¹¹ The yield was calculated on the basis of unrecovered ketone. C. Prepared from sodium nitrite and sulfuric acid.

١

o-O₂NC₆H₄CCOCO₂H o-Nitrobenzonitrile

# TABLE I-Continued

KETONES

Reference

Yield %

	236	237	i c	237	9	404	42	41	!	43		43	ç	238	239	239	240	38	241					
	61	]	ì	}	]	ı	56	Nearly	quant,	Nenrly	quant.	Nearly	quant.	8	1	1	09	ļ	ļ					
Products	n. Arul Alkul Monoketones-Continued	Benzil monoxime	2-Furoio acid	2-Furnidchyde oxime	2-Furoio acid	Benzaldehyde ozimo	2-0ximino-1-indanone	2-Oximino-1-indanono	2-0ximino-3-methyi-1-indanone		2-Oximino-5,6-methylenedioxy-1-indanono		2-Oximino-5,6-dimethoxy-1-indanome		a-Oximinonnrecin	a-Oximinonornarcein	a-Oximinomethyinyarhetii	Oximinomethyl-4-quinolyl Actoric	2-O ximino-1-(4-quinoly1)-1-propanono	o-02NG6H4CCOCO2H *	== ;	Z.	O (as bisphenylhydrazone)	
Method	B. Arul A.	MACONO NACCHA	Carring No.	Catalono, viii	ONO Ne	Camborne	, CMI	DE CHOOS	Chillonol HOI	Contillation	DH ONO HO	Chillone, acc	C,Huono, HC1		C.H.10NO, NaOC2Hs	C, HONO, NAOC, H	C, H, ONO, NAOC, Hs	C.H.ONO, NaOC2Hs	C,H,ONO, NaOC,Hs	NnNO2, CH3CO2H				
	Starting Compound		Deserventation (Cont'd)	Desoxyfuroln		Denoxybenzfuroin		1-Indanono		3-Methyl-1-indanone		5,6-Methylenedioxy-1-indanone		5,6-Dimethory-1-indanono		Narcein	Normarcein	Methylhydrastein	Methyl 4-quioolyl ketone	Ethyl 4-quinolyl ketooe	o-Nitrophenyipyruvic neig			

C45H10NO C55H10NO C55			C. p.Diketones 3-Oximino-2,4-pentanedione	44	52, 51 46
4-Oximino-1-phenyl-1,3-butanedione Ny03, ether Ny03, cyle (1900-1-phenyl-1,3-butanedione Ny03, ether Cycimino-1-phenyl-1,3-butanedione Ny03, ether Cycimino-1-o-methoxyphenyl-1,3-butanedione Ny03, ether Cyclinino-1-o-methoxyphenyl-1,3-butanedione Ny03, ether Cyclinino-1-o-methoxyphenyl-1,3-butanedione Cyclinino-1-o-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione Cyclinino-1-phenyl-1,3-propanedione Cyclinino-1-phenyl-1,3-propanedione Cyclinino-2-phenyl-1,3-cyclohexanedione Cyclinino-2-phenyl-1,3-cyclohexaned		C ₆ H ₁₁ ONO HNO ₂	3-Oximino-2,4-hexanicatione	86 43	242 46
HNO2  2-Oximino-1-o-methoxyphenyl-1,3-butanedione HNO2  2-Oximino-1-o-p-dimethoxyphenyl-1,3-butanedione Na ₂ O ₃ , ether C ₃ H ₁ 10NO, HCl 2-Nitroso-1,3-diphenyl-1,3-propanedione S ₂ Nitroso-1-phenyl-3-p-methoxyphenyl-1,3-propanedione C ₃ H ₁ 10NO, HCl 2-Oximino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione C ₃ H ₁ 10NO, HCl 2-Oximino-5-penyl-1,3-cyclohexanedione HNO2  CH ₃ ONO, NaOC ₂ H ₅ 2-Oximino-5-penyl-1,3-cyclohexanedione CH ₃ ONO, NaOC ₂ H ₅ 2-Oximino-5-penyl-1,3-cyclohexanedione CH ₃ ONO, NaOC ₂ H ₅ 2-Oximino-5-penyl-1,3-cyclohexanedione HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-minocyclohexane-3,5-dione HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-minocyclohexane-3,5-dione HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-pyrene HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-minocyclohexane-3,5-dione HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-pyrene HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-minocyclohexane-3,5-dione HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-pyrene HNO2  HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-pyrene HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-pyrene HNO2  HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxir-pyrene HNO2  1-(3-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi		C,H1,ONO, HCI N2O3, ether N2O3, C2H6OH, NaOC2H5 HNO3,	4-Oximino-3,5-heptaneduoue 2-Nitroso-1-phenyl-1,3-hutanedione 2-Oximino-1-phenyl-1,3-hutanedione 2-Oximino-1-phenyl-1,3-hutanedione 2-Oximino-1-phenyl-1,3-hutanedione		50 44 51 49
Ca, Hillono, HCI 2-Oximino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione chilono, HCI 2-Oximino-1-phenyl-3-p-mitrophenyl-1,3-propanedione chilono, HCI 2-Oximino-1,4-diphenyl-1,3-cyclohexanedione chilono, HCI 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione chilono, HCI 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione chilono, HCI 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione chilono, HNO2 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione chilono, HNO2 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione chilono chilon	1-o-Methoxyphenyl-1,3-butanedione 1-o,p-Dimethoxyphenyl-1,3-hutanedione Dihenzoylmethane	HNO2 HNO2 HNO3 NyO3, ether NyO3, ether NyO3, ether	2-Oximino-1-o-methoxyphenyl-1, 3-hutanedione 2-Oximino-1-o,p-dimethoxyphenyl-1,3-butanedione 2-Nitroso-1,3-diphenyl-1,3-propanedione 2-Nitroso-1,3-diphenyl-1,3-propanedione 2-Nitroso-1,3-henyl-3-p-methoxyphenyl-1,3-propanedione	50-60 80 35	50 48 50 50
c ₃ HilONO, HCl 2-Oximino-1-phenyl-3-p-nitrophenyl-1,3-propanentone   C ₃ HilONO, HCl 2-Oximino-1,4-diphenyl-1,3-butanedione   C ₃ HilONO, HCl 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione   C ₃ HilONO, HCl 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione   C ₃ HilONO, HCl 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione   C ₃ Hilono-5-phenyl-1,3-cyclohexanedione   C ₃ Hilono-5-	1-1,3-propane-	CsH110NO, HCI	2-Oximino-1-phenyl-3-p-methoxyphenyl-1,3-propanedione	1 1	50
CH50NO, NaOC2H5 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione HNO2 2-Oximino-5-phenyl-1,3-cyclohexanedione HNO2 2-Oximino-5-phenyl-1,3-cyclohexanedione HNO2 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oximinocyclohexane-3,5-dione 1-2,6-dir HNO2 HNO2 HNO2 A,9-Dinitroso-3,5-3,10-tetraketo-3,4,5,8,9,10-hexahydro-pyrene R, I m.N. I m. I m.N. I m. I m.N. I m.N. I m.N. I m. I m	I-Phenyl-3-p-nitrophenyl-1,3-propanedione 1,4-Diphenyl-1,3-hutanedione 5-5-Dimethyl-1,3-avvlobazanedione	C ₆ H ₁₁ ONO, HCl C ₆ H ₁₁ ONO, HCl HNO ₂	2-Oximino-1-phenyl-3-p-nitrophenyl-1,3-propaneurone   2-Oximino-1,4-diphenyl-1,3-butanedione 2-Oximino-5,5-dimethyl-1,3-cyclohexanedione	1 1 2	243
HNO2  2-Orimino-5-phenyl-1,3-ecyclohexaneuone  -, Quant, 2-Nitroso-1,3-indanedione  1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- minocyclohexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi- minocyclohexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi- minocyclohexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi- minocyclohexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi- minocyclohexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi- pyrene  1, G-Methoxy-4'-hydroxyphenyl)-2,6-dicarbethoxy-4-oxi- pyrene  R, Ind. Periodicarbethoxy-4'-acetoxyphenyl)-2,6-dicarbethoxy-4-oxi- pyrene  R, Ind. Derivatives (Amnono-Ketores)  R, Ind. Derivatives (Amnono-Ketores) 2-Caximinomethyl-3,3-direthylpseudoindole 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarbethoxy-4-oxi-  R, Ind. Derivatives (Amnono-Ketores)		CH3ONO, NaOC2Hs	2-Oximino-5,5-dimethyl-1,3-cyclohexanedione	1 1	46
1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi-  nuinocyclobexane-3,5-dione 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- nuinocyclobexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarhethoxy-4-oxi- nuinocyclobexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarhethoxy-4-oxi- nuinocyclobexane-3,5-dione 1-(3'-Methoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- sxahydro- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-hoxy-4'-acetoxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarhethoxy-4-oxi- 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dicarhethoxy-4	5-Phenyl-1,3-cyclohexanedione 1,3-Indanedione	HNO2 HNO2	2-Oximino-5-phenyl-1,3-cyclohexanedione 2-Nitroso-1,3-indanedione	-, Quant, (erude)	54, 53
mneoyetonezane-a, 3-duone 1-(3'-Methoxy-4'-oetoxyphenyl)-2,6-dicarbethoxy-4-oxi- 1-(3'-Methoxy-4'-oetoxyphenyl)-2,6-dicarbethoxy-4-oxi- 1-(3'-Methoxy-4'-oetoxyphenyl)-2,6-dicarbethoxy-4-oxi- minocyclobezane-3,5-dione minocyclo	1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-di-	HNO2	1-(3'-Methoxy-4'-hydroxyphenyl)-2,6-dicarhethoxy-4-oxi-		244
minocycloheznne-3,5-diona sxahydro- HNO2 4,9-Dinitroso-3,5,8,10-tetraketo-3,4,5,8,9,10-hexahydro- 89 pyrene R. Iralve Derivatives (Azmono-Kelones)  te HNO2 2-Oximinomethyl-3,3-dimethylpseudeindole NNO3 2-Oximinomethyl-3,3-diethylpseudeindole - 2-Oximinomethyl-3,3-diethylpseudeindole - 3-oximinomethyl-2-formoximeindoleninium 1-porchlorate - 66	carbethoxycyclohexane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-di-	HNO2	minocyclobezane-3,5-dione 1-(3'-Methoxy-4'-acetoxyphenyl)-2,6-dionbethoxy-4-oxi-	88	546
ie HNO2 2-Chiminomethyl-3,3-dimethylpseudeindole — 2-Chiminomethyl-3,3-dimethylpseudeindole — 2-Oniminomethyl-3,3-diethylpseudeindole — 2-Oniminomethyl-3,3-diethylpseudeindole — 2-Oniminomethyl-3,7-di	carhethoxycyclohexane-3,5-dione 3,5,8,10-Tetraketo-3,4,5,8,10-hexahydro-	HNO2	minocyclohexane-3,5-dione 4,9-Dinitroso-3,5,8,10-tetraketo-3,4,5,8,9,10-hexahydro- pyrene	83	56
**************************************	1000		crivities (Ammono-Kelones)		99
	2-Methyl-3,8-diethylysenkhinka 1-8,8-Trlinethyl-Smethylenedihydrolindsle	HNO. HNO. HNO. HCO.	2-Cammomethyl-3,3-dimethylpseudoindole 2-Oximinomethyl-3,3-diethylpseudoindole 1,3,3-Trimethyl-2-formoximeindoleninium 1-perchlorate	118	245 67

* These products were obtained when the reaction was run in hot aretic acid. No reaction took place at norm temperature. When an squeeus solution of e-nitro-Phenylpynuvic acid was boiled with excitum nitrite (2 equiv.) and Aprinx-Moric acid (2 equiv.), enterdenerating was produced in \$555 yield.

\( \lambda\) quantitative yield of the extresponding tribetone was obtained when a beneare solution of the Ekriste was trested with nitrons funce.

#### ORGANIC REACTIONS

Yield 97. Reference		75-80 246		ર્હે *	052		-, 60 * 70, 252				50 1 202, 253, 254			257	6	60 259, 33			_		02					2 2	•	202, 202	263	262	264	50.51 75,72	73	20-30 263	
eta-Keto Acids, Esters, and Related Compounds	Products	A. B-Keto Acids	Oximinoncetooo	Oximinoacetono	Oximinoacotono	1-Oximino-3-ethoxy-2-propanone	Diacetyl monoximo	Dincetyl monoxime	Diocetyl monoximo	3-Oximino-2-pentacone	3-Oximino-5-ethoxy-2-pentanone	3-Oximino-2-hoxBnone	3-Oximino-4-methyl-2-pentanone	3-Oximino-5-boxene-2-one	3-Oximino-5-mothyl-2-hoxanone	3-Oximino-2-ootanons	3-Oximino-6-mothyl-2-beptanone	3-Oximino-4-methyl-2-decanone	3-Oximino-4-phenyl-2-butanone	3-Oximino-4-phonyl-2-hutanone	3-Oximino-4-m-tolyl-2-butanone	3-Oximino-4-beptanone	5-Orimino-4-octanone	3-Oximino-2-methyl-4-heptanone	2-Oximino-6-methyl-4-beptanone	2-Oximino-3-octanono	3-Oximino-4-nonanone	-Oximin onno nionhenone		A-Oximinobutyrophonous	p-Oximino-p-knowaria	7-Oximino-5-ketobexanoto actd	Dioximinoacetone	Dioximinoacetone	
$\beta$ -Keto Acids, 1		Method	ONLI	HOIL HAO	HOIL HAO	HOIL HAOT	IOUI; IIOU	IINO3	HOII; HNO		HOH; HAO!	HOII; HNO	HOH; HOO!	HOII; HNO3	HOH; HOG	HOLE HOS	HOH; HNO3	HOH; HNO	NOCI	HNO2	HOII; HNO	HOH; HOG	HOH; NOC!	HOH: NOC	HOH: NOCI	HOH; NOCI	NOC!	NOCI	HOH; HNO2	HOH; HNO2	HOIL; HNO2	HOH: HNO2	HNO2	HNO,	
		Starting Compound		Pilly   acresaretate			Ethyl z-ethoryacetoacetato	nation o-methylacetonectate	Fring amethylacetonectato		Eshal agthylacetoacetato	Pithyl o-2-thoryethylacotoncetate	Ethyl or propylaretoacetato	Ethyl o-inopropylacetoncetato	Ethyl a-ullylacetoncetate	Ethyl o-isobutylacetoncetate	Ethyl or anylacetoncetate	Fthyl o-isoamylacetoscolate	gere-Octylaeotoacetic acid	g-Bensylncetonectioneid	Ethyl a-benaylacetoncetate	Ethyl a-m-xylylacetoncetate	Ethyl o-ethyl-8-ketocaproato	Ethyl a-propyl-g-ketocaproato	Ethyl a-isopropyl-g-keto eaproato	Ethyl 2-ethyl-3-keto-5-mothylhexanoate	a-Methyl-B-ketocaprylio acid	a-Ethyl-g-ketoenprylie neid	Eshul _methylhenzovlacetate	Ethyl gethylbengoylacetate	Dietal acetulanceinste	Dietay neety succession	Diotaly denotylkanama Ambandionshowelia acid	שולית וורוויות בביל ייי יייי	

Diethyl a,a'-diacetylsuccinato (Con'd) Diethyl a-acetyl-a-methylsuccinate 2-Carboxycycloberanone	нон; нио; нон; пио; пио;	3-Oximino-2,5-hexaneliono 3-Aeryl-1-anethyl-5-isoxatolous (?) 1,2-Cyclohexanelione monoximo 2,4-Dinimino-cyclohexanono	Very small	888 1
2-Carbethoxycycloliexanone	HOH; HNO;	1,2-Cyclohazandinno monorimo 1,2-Cyclohezandiono monorimo	; ? ? ?   !	. V. 9.
2-Carbethoxy-5-methyleyclohexanone 2-Carbethoxy-6-methyleyclohexanone Menthonecarboxylio acid	HOH, HNO; HOH; HNO; HNO;	2-Oumno-5-methyleycloberthone 2-Oumno-5-methyleycloberthone 4-Oumnomenthone	. <del></del> 1	. K. E.
	II. A	H. p. Keta Externach Amilia	d American	1.5%
Methyl acetoacetato	N0;50,111	Methyl oximinoarelate Edwl oximinoarelate	;	27.4, 25.2
Ethyl acetoacetata	IINO;	Ethyl a-oximina acetuacetato	•	13. (1) 15. (1) 14. (1)
	OWI.	Selection of the select	(E) (C)	5, 14, 150,
	11302			1 23, 270,
			6604 300	100
	IINO;	Ethyl a-oximinoacetorectate		
	CII,0NO, IICI	Ethyl a-oximinoacetoaretale	e de	* 1
	CII,ONO, NAOC, II,	Ethyl a oximinoscetoscetste	<b>:</b> .	
	Csilitono	Ethyl a-aximinoacetoacetate	ţ	T 1
Isobutyi acetoacetate	NO,SO,III	Isobutyl a-otiminoscetate	ę.	i i
Ethyl &-ketovalerate	HNO	Ethyl a oximino paketovalerate	Ļ	3, 3
Methyl benzoylacetate	HNO:	Methyl a-oximinobenxoylacetate	2	***
Ethyl benzoylaeetate	HNO:	Ethyl a-nximinabensoylacetate	i. 9	
	CsHriono	Ethyl a oximinobensoylacetate	í	7
Methyl o-methoxybenzoylacetate	HNO;	Methyl agriming-methoryhears) lacitic		e e
Methyl m-methoxybenzoylacetata	HNOz	Methyl a-oximina-m-metharylwaroylacetate	2	
Methyl p-methoxybenroylacetate	IINO;	Methyl a-oximino-p-methoxybenzoybacetale		
Ethyl m-methoxybenzoylacetate	HNO	Ethyl a oximino - methoxybenxoylacetein	;	7.8
Ethyl p-methoxybenzoylacetate	HNO;	Ethyl acomining-penetherybenenylacetate	:	7 7
Ethyl p-nitrobenzoylacetato	N:O3, ether	Ethyl amenina-p-nitrobenzoylacetste	į	80
Note: References 191-316 are listed on pp. 375-377.  * The yield was based on the dioxime industral	pp. 375-377.			

* the yield was based on the dioxime isolated.

† The yield was calculated on the basis of unrecovered eater.

† The yield was based on the diketone isolated.

8

# TABLE II-Continued

	Harsh sons some	STATES ACIDS ESTERS, AND RELATED COMPOUNDS	Yield	
	p-Meio Meio	The state of the s	%	Reference
	Mathod	ronner		
Starting Compound	refull control	n o rear Estore and Amides-Continued	Almost	96
	Calliono (1 ed.), 11Cl	Diethyl a-oximino-3-ketoglutarato	quant.	į
Diethy I acetonedicarboxydate			ļ	76
	Calinono (1 eq.), IICI	Diethyl a-oximino-p-ketoginum.	99	/6
	CallinoNo (3 eq.), HCl	3,5-Dicarbethoxy-i-nyaroxyisoxario	i	787
	HNO	Ethyl a-oximinoluroyinceuro	65	1/1
Ethyl (uroylacetate	INO.	Ethyl a-oximinopicolinoyincounto	i	172
Ethyl picolinoylacetato	INO	Ethyl a-oximinonicotinoylneethu	Į	283
Ethyl nicotinoy lacetate	-CNII	a-Oximinotetronio acid	ļ	284
Tetronio neid	INO	a-Oximino-y-pheayltetronie neiu	i	285
Phenyltetronic acid	INO	a-Oximinobenzotetronic neid	i	286
Benrotetronia acid	HNO	a-Oximinonectoacetanilide	i	287
Acetoacetanilido	TOWN STOCK	a-Oximinoacctoncetanilido	ı	287
	000	a-Oximinoacctoacet-o-toluidide	i	287
.Artoncet-o-toluidide	502	a-Oximinoncetoncet-p-toluidide	i	287
Aretoncet-p-toluidide	500	a-Oximinoacetoacet-2,4-dimethylanilido	. 1	287
Acetoacet-2, f-dimethylanilide	100 K	a-Oximino-N-a-naplithylacetoneetamide	1 !	287
N-a-Naphthylacetoacetamide	1000	a-Oximino-N-8-naphthylacetonectamide	ä	82
N.g. Naphthylacetoneetamide	1002	Ethyl a-nitrosopropionato	3	68
Fillyl a-formylpropionate	N ₂ O ₃	Delay oriming-?-nitrophenylacetate	<b>∵</b>	700
return a formylation via cetato	N203	Littly Committee and Secondary	29	288
New Alanda - market also soften potats	NO ₂ SO ₂ H	Methyl a-daminopropries	l	10
Methyl dinemy meetoneers	NOCI	Methyl a-oximinopropionate	Į	208
of of occupant of the state of	NO,SO,H	Ethyl a-oximinopropionate	Ouant.	82
Ethyl a-methylndetoncedia	0.N	Ethyl &-nitrosopropionate	32	89
	ONH	Ethyl a-oximiaopropionate	1	68, 289
	HOH: HOH	a-Oximinopropionic neid	8	92
	C.H.ONO. 85% H.SO4	Ethyl a-oximinopropionate	3	268, 288, 29
open and the state of the state	HrOS*ON	Ethyl a-oximinobutyrate	2 4	855
Ethyl a-ethylacelouectate	0,0	Ethyl a-nitrosobutyrate	·	291
	HOH: HOH	a-Oximinobutyrie acid	8	200
	C,H,ONO, 85% H,SO4; HOH	a-Oximinobutyrie acid	8	268, 288
Ethyl a-n-propylacetoacetate	HtOSON	Ethyl a-oximinovalerate	3 1	292
	H002; H0H	a-Oximinovalene ucio		

NITROSAT	ION U	)F ALIPHAL	10 02	Hibor M-	. • • • • • • • • • • • • • • • • • • •
92 290, 288 288 164 85 290 162 9, 92 290, 288 92 91	288 259 288	102 155 293 88 92 92 84, 85	118 92 85	294 9, 92 9	9 9 260a 92 101
85 -, 75 93 74 Quant. 84 86, 89 -, 90 70 70 70 70	818,	ğ     1 %   8	861	70-80 85, 89 45 Small	Small 52 62 Small 87 62
α-Oximinovaleric acid Jaohutyl α-oximinovalerate Ethyl α-oximinoisovalerate Ethyl α-oximinoisovalerate Ethyl α-oximinoisovalerate Ethyl α-oximinocaproate Ethyl α-oximinocaproate Chyl α-oximinocaproate Chyl α-oximinocaproate α-Oximinocaproa acid α-Oximino-β-methylvalerio acid α-Oximino-β-methylvalerio acid Ethyl α-oximino-β-methylvalerio	Ethyl 2-nitroso-3-meulylickinouv Ethyl 2-oximino-5-methylhexanoate Ethyl 2-oximino-5-methylhexanoate Tethyl a-aximino-8-methylpelargoato	Linyl a-oximino-p-incomputate Ethyl a-oximino-p-incomputate Ethyl a-oximinopropionato Ethyl a-oximinopropionato Ethyl a-oximinopropionate Diethyl oximinoprutyrate Diethyl oximinosuccinate	Diethyl a-oximinoglutarate Diethyl a-oximinoglutarate Diethyl a-oximinoglutarate	Diehyl α-nitroso-α-acetylsuccinide Ethyl α-oximino-β-phenylpropionate α-Oximino-β-phenylpropionic acid α-Oximino-β-phenylpropionic acid α-Oximino-β-phenylpropionio acid	
C,H ₂ ONO, 85% H ₂ SO,; HOH NO ₂ SO ₃ H NO ₂ SO ₃ H C ₂ H ₂ ONO, Na ₂ O ₂ H ₅ C ₂ H ₃ ONO, Na ₂ O ₂ H ₅ C ₂ H ₃ ONO, 85% H ₂ SO,; HOH NO ₂ SO ₃ H NO ₂ SO ₃ H NO ₂ SO ₃ H	N ₂ O ₃ NO ₂ SO ₃ H C ₂ H ₅ ONO, NaOC ₂ H ₅	NO;SO;H NO;SO;H NO;SO;H No;0, No;OzH; NO;SO;H C,H;ONO, 85% H;SO; N;O;	NO2SO3H C2H6ONO, KOC2H6 C4H3ONO, 85% H2SO4	N ₂ O ₃ NO ₂ SO ₃ H C4H ₂ ONO, 85% H ₂ SO ₄ ; HOH C ₄ H ₂ ONO, 85% H ₂ SO ₄ :H ₂ PO ₄ (1:2); HOH C ₄ H ₂ ONO, (CH ₂ CO) ₂ O;	HOH C.H.ONO, HCO.H. HOH C.H.ONO, HCI, HOH C.H.ONO, NnOC.H.; HOH HNO.; HOH C.H.ONO, 85% H.SO.; HOH -C.H.ONO, NnOC.H.;
Ethyl a-n-propylacetoacetate (Cont'd) Isobutyl a-n-propylacetoacetate Ethyl a-isopropylacetoacetate Ethyl a-n-butylacetoacetate Ethyl a-isobutylacetoacetate	Ethyl æisonnylacetoacetate	Ethyl a-sec-octylacetoacetato Ethyl a-2-bromoethylacetoacetate Ethyl a-3-diethylaminopropylacetoacetate Ethyl a-methyl-f-ketovalcrato Ethyl a-ethyl-f-ketovalcrato	Dicthyl a-acetylglutarate	Diethyl α,α'-diacetylsuccinato Ethyl α-benzylacetoacctato	Ethyl m-xylylacetoacetate Ethyl a-p-methoxybenzylacetoacetate Ethyl 3,4-methylenciloxybenzylacetoace-

Note: References 191-316 are listed on pp. 375-377.

# TABLE II—Continued

	B-Kero Acids, Es	B-Keto Acids, Esters, and Related Compounds	Yield	Reference
,	Method	Products	<b>o</b> /	
Starting Compound	B. A-Keto 1	B. g-Kelo Esters and Amides—Continued	l	2 9
Methyl æmethylbenzoylacetate	), (NaOC)	Methyl œ-oximinopropionave Methyl œ-oximinopropionave Ethyl œ-nitro30-œ-methylbenzoylacetave	Quant.	295 82
Ethyl & methylhenzoylacetate Diethyl henzoyleuccinate 2-Carhethorycyclopentanone	N2O1 N2O1 C2H4ONO, NaOC2H8 C2H6ONO, CH3COCI	Diethyl oximinosuccinate Diethyl oximinosuccinate 2. Nitroso-2-oxibethoxy eyelopentanone	8 <del>6</del> 1 8	8 84 84
2-Carhethoxycyclohexanone	C2H5ONO, NAOC2H5 C2H5ONO, CH3COCI	Diethyl g-oximinopimica- 2-Nitroso-2-carbehaxycyclohexanone riseki aoximinomethyladipate	25-30	88
2-Carbethoxy-4-methylcyclopentanone	N2O3, NaOC2H3 C2H5ONO, NaOC2H6 C3H5ONO, CH2COC1	Dictary a comming of the property of the prope	30 25-51	38 8 8 36 8 8
2,6-Dicarbethoxy-1,4-cyclohexancdione c-Acetyl-y-butyrolactone	N.O., ether HNO. C.H.ONO, HCI	2,5-Dinitroso-2,5-dion beliaby-1,3-5, according to the continuory-butyrolactone a-Oximinory-butyrolactone a-Oximinory-butyrolactone a-Oximinory-a-fallerolactone	70 85-91 81	297 95 94 95
α-Acctyl-&chloro-γ-valerolactone	HNO ₂ NO ₂ SO ₃ H	a-Oximino-b-chloro-y-valerolactone	8 8	888
a-Methyltetronic scid	NaNO2 N2O3, CH3CO2H	a Nitroso antendronio acid	57 65	100
-Ethyltetronic acid -Benzyltetronic acid Ethyl N-methyl-N-phenylcarbamylpyru-	NaNO2 NaNO2 C ₆ H11ONO, NaOC2H5	a-Oximinoced one asset a-Oximino-6-phenylpropiony)glycolic acid Ethyl oximino-(N-methyl-N-phenylcarbamyl)pyruvate	1 52	100 46
vate	C. p-Imino Acids	C. p-Imino Acids and Esters and p-Keto Imino Ethers	١	81
Ethyl <i>β-</i> aminecrotonate	HNO2 C ₅ H ₁₁ ONO	Ethyl α-oximino-β-iminobutyrate Ethyl α-oximino-β-intosiminobutyrate (ammonium salt) Ethyl α-oximino-β-intosiminobutyrate (ammonium salt)	1 1	90 40
Monoethyl a-cyano-β-iminoglutarie acid 9-Phenyliminobutyric acid Benzoylacetimido ethyl ether	HNO2 HNO2 CSH11ONO, NH3 CSH11ONO, HCI HNO2	Ethyl arcylinnopropionaldehyde oxime Orhinnopenzoylacetamidine Oximinobenzoylacetamidine Oximinobenzoylacetate Ethyl aroximinobenzoylacetate	1   1   1	81 26 26 26
Benzoylacctimido etbyl etber hydrochloride NaNO2 Note: References 191–316 are listed on pp. 375–377.	NaNO ₂ 375-377.	Oximinobenzoylacetimido cthyl ether		

## TABLE III

,	Reference	6	6	102	6	12	102	12	12	ტ	6	101	297	298	103	104, 107, 108, 113	104	105	106		111	103, 110	112	299		6	∞ .	6	∞ •	297	297	
Yield	%	5	2 62	70-80	84	75-85	62	40-48	8	8	26	85-90	34	i	i	- '02' '09	# 06 <del>-</del> 08	50 (max.)	Consider-	nble amt.	90-95	85-90	87	Consider-	able amt.	65	8	70	35 1	7.5 50-56	1	
Malonic Acids, Esters, and Amides	***************************************	Products	a-0 ximinopropionic acid	a-Oximinobutyrie neid	a-Oximino-r-butyrolaotono	a-Oximinocaproic acid	a-Oximinoisoenproio neid	a-Oximino-8-phenylpropionic acid	a-Oximino-8-phenylpropionic acid	α-Oximino-β-plienylpropionic acid	α-Oximino-β-phenylpropionic neid	$\alpha$ -Oximino- $\beta$ -p-methoxyphenylprophonic ucid	g-Oximino-6-5,4-metaylenedibay pacity in the	. O.: 2 at the limit of a cetyl-2-propanone	L-Oximino-3-primaring cocos - Francis	Umethyl oximinomalonave Diethyl oximinomalonate	•	Diethyl oximinomalonate	District of the majorate	Dietnyi oximinomalonine		Diediyi oximinorinabilako	Diethyl oximinomalonate	Diethyl oximinomalonate	Ethyl a-oximinopropionate		G-Caminobudy its action	Chiningentrio neid	Ethyl a-oximino-8-phenylpropionate	a-Oximino-8-phenylpropionio neid	a-Nitroso-a-carbethoxy-y-butyrolactono a-Nitroso-a-carbethoxy-y-butyrolactono	
Malonic Acros,		Method	IDH ONO HO	Chigono, no:	No.NO. *	C.H.ONO. HCI	C.H.ONO. HCL +	No.NO.	HWO.	C.n.ONO. HCl	C.H.ONO. HCI	C,H,ONO, HCI	4-C311,0NO, HC1	HNO	HOH, HNO2	CII,ONO, NaOCH, HNO2		HNO,	N2O3, NaOC2H5	N ₂ O ₃ , N _n OC ₂ II ₅		N2O3, NaOC2HS	CH10NO, NaOC1116	C,H,ONO, NaOC,H,	N ₁ O ₁ , N _n OC ₁ II ₅	HOH H DO M ONO M D	CAHIONO, NAOCIAS, MOR	CHISONO, NEOCHHS	Call ONO NaOCalls	C,H,ONO, NaOC,HS; HOH	HNO2 N-O3	
			Starting Compound	Methylmulonic acid	Ethylmalonic acid	g-Bromoethylmalonic acid	n-Butylmalonic acid	Isobutylinalonic acid	Benaylmalonic acid			. Neathern Charles and a side	3.4. Methylenedioxybenzylmalonic acid	-Carbory butyrolactone	Diethyl phthalimidoacetylmalonath	Dimetlyl malonate	Diethyl inniburate								Diethyl methylmalonate		Diethyl ethylmalonate	Diethyl n-butylmalonate	Distral Languages	The transfer of the second of	a-Carbethoxy-y-butyrolactona	

• The reaction versel was sealed tightly before it was shaken with each portion of the reagent. † No traction took place with sodium nitrite and sulfuric acid.

Note: Beforences 191-316 nte listed on pp. 375-377.

There makes of nitrous acid were used in this experiment. One malo and two males gave only 50 and 63% yields, respectively, of oxime.

# TABLE III-Continued

Yield Befarence	70 158, 114 70-83 157 94 8 02 300	301	70 115 40 116 Quant. 117	Quant. 117 Quant. 117 Quant. 117 Quant. 117 — 117 Quant. 117 Poor 117 ride — 115	which happens and the state of the second
Malonic Actos, Esters, and Amdes	Products Ethyl a-oximino-b-cynnovalerate Ethyl a-oximino-b-cynnovalerate Ethyl a-oximino-b-diethylaminovalerate	Diethyl 1-(o-nitrophenyl)-3-oximinokuusuusu	Ethyl a-oximinoacetoaceiato Oximinomalonamido Oximinomalonamide Oximinomalonamide	a-Oximino-N,N'-dimethylmalonamide a-Oximino-N,N'-diphenylmalonamide a-Oximino-N,N'-diphenylmalonamide a-Oximino-N,N'-dip-o-tolylmalonamide a-Oximino-N,N'-dip-p-tolylmalonamide Ethyl a-oximino-N,N'-p-tolylmalonamide a-Oximino-N,N'-a-naphthylmalonamide a-Oximino-N,N'-dip-naphthylmalonamide a-Oximino-N,N'-diphenylmalonamide o-Oximino-N,N'-dimethyl-N,N'-diphenylmalonamide o-Oximino-N,N'-dimethyl-N,N'-diphenylmalonamide oximinomalonyldiurethane	
Malonic A	Method C:H;ONO, NnOC:H; C:H;ONO, NnOC:H;	Callsono, NaoGalis	11NO3 11NO3 N3O3, 113O	NOCI NOCI NOCI NOCI NOCI NOCI NOCI NOCI	275_277
	Starting Compound Dietigly eyanopropylmalonate	Diethyl 3-diethylaminopropylmalomate Triethyl 1-(0-nltrophenyl)propano-1,3,3-	tricarboxylate Dichyl acetylmalonate Malonanide	N,N-Dimethylmalonamide N,N-Diplorylmalonamide N,N-Di-e-tolylmalonamide N,N-Di-p-tolylmalonamide N,N-Di-p-tolylmalonamide Ethyl N-p-tolylmalonamide N,N-Di-g-naphthylmalonamide N,N-Di-g-naphthylmalonamide N,N-Di-g-naphthylmalonamide N,N-Di-g-naphthylmalonamide N,N-Di-g-naphthylmalonamide N,N-Di-g-naphthylmalonamide	775-777 and - 11-1-1 - 375-377

Note: References 191–316 are listed on pp. 375–377. f The salvent is noted because poor yields were obtained with chloroform as solvent and when the nitrosation was run with liquid nitrosyl chloride in a sealed tubo

## TABLE IV

ARYLACETIC ACIDS AND ESTERS

	Reference	118	119.	16	120	120	
Yield	% {	Good	1	8	3 1	1	
ARYLACETIC ACIDS AND ESTERS	Products	Ethyl a-oximinophenylacetate Ethyl a-oximino-p-bromophenylacetate	Ethyl a-oximino-o-nitrophenylacetate	Ethyl a-oximino-p-nitrophenylacetate	3-Carbomethoxy-6-nitrobenzisoxazole	3-Nitro-4-Ommino-anitrophenylacetate	
ARYLACE	Method	Clesono, Kocius	C.H.,ONO. NoOC,Hs	CsH110NO, NaOC2H5	CsH110NO, NAOCH3	Cshnono, HCl	Cshilono, noi
	Starting Compound	Ethyl phenylacetato	Ethyl p-bromophenylacetate	Ethyl o-nitrophenymeetho	Methyl 2,4-dinitrophenylacetate	2-Nitro-f-aminophenylacetic acid	Ethyl 2-nitro-f-aminaphenylacetate

Note: References 191-316 are listed on pp. 375-377.

### TABLE V

Nitriles

							~	_			' '		-			_			-	`	,,,,		4)(	,11	71.	1.	/11/L	,			071
Reference	123	122	123	122, 123, 125	124, 302	123	122	129, 122	129	122	126	126	126	303	128	127	127	128	128	128	133	132		132		132	118	131	130	304. 305	
Yield %	90-95	j	Poor	87~100	j	Poor	70	8,	1	ļ	ļ	Small	J	ţ	20	J	J	55	61	63	ļ	j	I	ļ	ļ	Very small	amount Small	j	88-92	99	
Products	Methyl oximinocyanoacetate	Methyl oximinocyanoacetate	Methyl oximinocyanoacetate	Ethyl oximinocyanoacetate	Ethyl oximinocyanoscetate	Ethyl oximinocyanoacetate	Oximinocyanoacetamide	Oximinocyanoacetylurea	N-Methyl-N'-oximinooyanoacetylurea	Ethyl oximinocyanoacetylcarbamate	a-0ximinobutyronitrile	Oximinophenylacetonitrile	a-Oximino-8-phenylpropionitrile	Oximinophenylacetonitrile	Oximinophenylacetonitrile	Oximinophenylacetonitrile	Oximino-p-bromophenylacetonitrile	Oximino-o-chlorophenylacetonitrile	Oximino-p-chlorophenylacetonitrile	Oximino-p-nitrophenylacetonitrile	α-Oximino-β-nitrosimino-β-phenylpropionitrile	a-Oximino-8-nitrosimino-8-phenylpropionitrile	Oximinobenzoylacetonitrile	a commody-hittosimino- $\beta$ -p-tolyipropionitrile (amnio-nium salt)	Oximino-p-toluylacetonitrile	α-Oximino-β-nitrosiminobutyronitrile (ammonium salt)	Dioximinosuccinonitrile	Oximinomalononitrile	a-Oximino-b-amino-p-ethoxy-p-hydroxypropionitrile	CH10,—CH-CH-CH1	CH2 CH2CONHC-0
Method	HNO,	HNO ₂	C,H110NO, NaOCH1	HNO ₂	HNO ₂	Chilono, Naoch	HNO ₂	NaNO ₂	HNO ₂	HNO ₂	RONO, KOC2H5	RONO, KOC2H5	Cefuono, Kochi	N2O3	RONO, NaOC2H5	Carriono, NaoCaris	Cennono, NaoCalle	CSHIIONO, NAOC2Hs	CHILDNO, NAOCH	HNO. TO	CH. ONO	Ospiloso	C,H1,ONO		C.HONO	Carriera	C,H110NO, KOC,H5	CHIONO NAOCAE	STITE OF THE STITE	C5H11ONO, NaOC2H5	
Starting Compound	Methyl cyanoacctute			Ethyl cyanoacctate			Cyanoacetamide	Cyanoacetylurca	N-Methyl-N'-cyanoacetylurca	Ethyl cyanoncetylcarbanate	Ethyl a-eyanobutyrate	Ethyl cyanophenylacetate	Phonological Company Proposite	z neuglacetomuna		n-Bromophenile antonitalle	9-Chlorophenylacetonitrile	p-Chlorophenvlacetonitele	p-Nitrophenylacetonitrila	8-Imino-8-phenylpronienitrile			heta-Imino- $ heta$ - $p$ -tolylpropionitrilo		\$-Iminobutyronitrile		Succinonitrilo Malononitrilo			Cynnedihydrocarvono	Note: References 101-316 and 11-213

## TABLE VI

	Reference	308, 307,	901	310	311	300	134, 310	#	110	312		186	314	315	137		197	: 1	137	1100, 121	111 601	90	2 9	140	140	140				Reference	145 146		143
i	Refer	1, 500 30	1, 306	309, 310	136, 311	2	134,	134	2, 310	. 8.	313	18	. 5				5 2	2 5	13 6	1001	1001	· •		1	1	2				Refo			
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	5																					xime			•	acid				ន			
	Products													olo.	શુ	ncid			neid	жіте	xime	2-Nitro-1-methylbenzaldchyde oxime		xime	14. Dinitrobenzophenono oximo	Phenyl-p-nitrobenzohydroximie acid				Products	üme		ево
:D3		olie acid		ic acid	ie acid	io acid	llo acid	lie acid	donitrole	lonitrolo	io acid	onitrolo	rolic acid	Cyclohexyl [Seudonitrole	Camphene recudonitrole	Hydroxyethyl nitrolio acid	fio acid	o neid	Hydroxyethyl nitrolio neid	o-Nitrobenzaldehydo oximo	p-Nitrobenzaldehyde oxime	thylbenzo	fonitrife	P.Nitronce tophenone oxime	cuzophen	obenzohy			••		Di-n-propyl ketone oxime		Oximinocyclopentadieno
OVEOUS		se et al miteralis acid		Ethyl nitrolic acid	Ethyl nitrolic acid	thy! nitrolle acid	'ropyl nitrollo acid	Propyl nitrolic acid	ropyl p-eudonitrole	ropyl perudonitrolo	Intyl nitrolie acid	Butyl preudonitrolo	sobutyl nitrolic acid	clohexyl 1	nphene p	droxyeth	Methyl nitrolio acid	Ethyl nitrolio acid	droxyethy	itrobenza	litrobenza	itro-f-me	4-Nitrogalicylonitrile	itroncetol	Dinitrob	nyl-p-nit	E. VII	1	ARBONS		Di-n-propyl ketone o	· Curantan	iminocyc
NITHO COMPOUNDS		:	7.	Ē	Œ	ä	-	1.1	ų	1.	Ē	n.	Į.	Ċ	٠ ق	II.	7	2	115	Z.	E	Z,	Z-	2	7	, ă	TARLE VII	7777	Нурносапромя		ä	4	ő
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		11.	•		**	-	**	13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03 13.03	<u>.</u>	•	_									navor Guldovo NaoCalla	ALCON CACHE	CHICAGO NAOCHIA	LONO NOCH	CALLONO MACCALL		CHILONO, NACCHIA				Method	NOCI, sunlight	NOCI, Minugue	CHIONO, NAOCHI
			H.S.C.		22	ioni.																				011150 021110	.375-377				NOCI,	100	C;1140
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												State Clair	A. Milly, buttere	markaniferration of the	Water-yelch exam	P. N. 13 P. A. S. P. S.	2.Nitra-Irelbaned		Witten frontenne	Skino-1, J. propanedio	- Mitrefeluene	P.N.trofeluene	Nitro-Paylene	2,4-Dinitroteluena	a Nitroethyfbensens	N.4. Dinitrodiphenylmethans	Nete: References 191-316 are listed on pp. 375-377.			,		£17.	Cythywntadiene Cyll(O)
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    252 Müller and Pechmann, Ber., 22, 2127 (1889).
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#### CHAPTER 7

### EPOXIDATION AND HYDROXYLATION OF ETHYLENIC COMPOUNDS WITH ORGANIC PERACIDS

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#### INTRODUCTION

Oxiranes ( $\alpha$ -epoxy compounds) and  $\alpha$ -glycols can be prepared from olefins by a variety of methods. One of the most important and most generally applicable of these is the oxidation of ethylenic compounds with organic peracids, as exemplified by the accompanying equations.

Depending upon the peracid employed and/or the operating conditions, either an oxirane  1,2,3  or an  $\alpha$ -glycol  2,4  can be obtained in good yield. Ordinarily the oxirane isolated can be hydrolyzed to the  $\alpha$ -glycol.⁵ It is important to note that the oxidation step both in epoxidation and hydroxylation reactions with organic peracids is the conversion of the olefin to the oxirane.

The literature on the epoxidation and hydroxylation of compounds containing an isolated ethylenic linkage is so extensive that no attempt has been made to include conjugated systems in a comprehensive fashion. However, occasional comments on  $\alpha,\beta$ -unsaturated acids are found on pp. 385 and 388, the preferential epoxidation of one ethylenic linkage in isoprene is described on p. 397, and a limited number of conjugated dienes and  $\alpha,\beta$ -unsaturated acids are included in Table I.

¹ Findley, Swern, and Scanlan, J. Am. Chem. Soc., 67, 412 (1945).

² Swern, Billen, and Scanlan, J. Am. Chem. Soc., 68, 1504 (1946).

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⁶ Swern, J. Am. Chem. Soc., 70, 1235 (1948).

#### SCOPE

#### Epoxidation

Perbenzoic Acid. The discovery that oxiranes can be prepared from ethylenic compounds by epoxidation with an organic peracid is generally credited to the Russian chemist, Prileschajew,6-9 who showed that perbenzoic acid is an efficient oxidizing agent for the epoxidation of isolated double bonds. This reaction is excellent for preparative pur-

$$\begin{array}{c|c} -C = C - + C_6 H_5 CO_3 H \xrightarrow{Organic} -C - + C_6 H_5 CO_2 H \\ \hline \end{array}$$

poses. It proceeds under mild conditions, and it is generally conducted in a non-reactive organic solvent, such as chloroform, ether, benzene, acetone or dioxane. The reaction time is usually short, but it varies with the number and nature of the groups attached to the ethylenic system.10 As a rule the yields are high.

Most investigators have preferred to prepare a solution of perbenzoic acid 3,11-15 for epoxidation. However, since perbenzoic acid can be prepared conveniently by the oxidation of benzaldehyde with oxygen,3,16-10 some investigators have treated solutions of benzaldehyde and the unsaturated compound with air or oxygen, the perbenzoic acid being consumed as it is formed. This application of the perbenzoic acid epoxidation technique, in which separate preparation and isolation of the peracid is avoided, has been applied to the oxidation of methyl oleate, 20 oleyl alcohol, 20 octenes, 21 oleic acid, 3.22 stilbene, 22 styrene, 22 and squalene,22 and good yields of oxiranes were generally obtained. When

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<sup>6</sup> Prileschajew, Ber., 42, 4811 (1909).
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⁷ Prileschajew, J. Russ. Phys. Chem. Soc., 42, 1387 (1910) [J. Chem. Soc. Abstr., 100, I, 255 (1910)].

⁸ Prileschajew, J. Russ. Phys. Chem. Soc., 43, 609 (1911) [C. A., 6, 348 (1912)].

⁹ Prileschajew, J. Russ. Phys. Chem. Soc., 44, 613 (1912) [C. A., 6, 2407 (1912)].

¹⁰ Swern, J. Am. Chem. Soc., 69, 1692 (1947).

¹¹ Braun, Org. Syntheses, Coll. Vol. 1, 431, 2nd ed. (1941).

¹² Hibbert and Burt, J. Am. Chem. Soc., 47, 2240 (1925). 13 Kolthoff, Lee, and Mairs, J. Polymer Sci., 2, 199 (1947).

¹⁴ Levy and Lagrave, Bull. soc. chim. France, [4] 37, 1597 (1925).

¹⁵ Tiffeneau, Org. Syntheses, 8, 30 (1928).

¹⁶ Jorissen and van der Beek, Rec. trav. chim., 45, 245 (1926). 17 Jorissen and van der Beek, Rec. trav. chim., 46, 42 (1927).

¹⁸ Jorissen and van der Beek, Rec. trav. chim., 49, 138 (1930).

¹⁹ van der Beek, Rec. trav. chim., 47, 286 (1928).

²⁰ Swern and Findley, J. Am. Chem. Soc., 72, 4315 (1950).

²¹ Pigulevskii, J. Gen. Chem. (U.S.S.R.), 4, 616 (1934) [C. A., 29, 2145 (1935)]. 22 Raymond, J. chim. phys., 28, 480 (1931).

aliphatic aldehydes, such as acetaldehyde and butyraldehyde, are employed instead of benzaldehyde, poor yields of oxiranes result.20,21,23

Epoxidation with perbenzoic acid has been employed in the preparation of oxiranes from an extremely large number and wide variety of ethylenic compounds (see Table I).

Monoperphthalic Acid. Another reagent that has been employed in the preparation of oxiranes is monoperphthalic acid; but this reagent, although efficient, has not been studied so extensively as perbenzoic acid, primarily because it offers only minor advantages in most reactions. When the epoxidation requires a long period of time for completion, however, the greater stability of monoperphthalic acid,24,25 compared to perbenzoic acid, is an advantage. Furthermore, since epoxidations with monoperphthalic acid are usually conducted in chloroform solution and the phthalic acid formed is insoluble, it is readily separated from the oxidation product. Although Böhme 26,27 was apparently the first to demonstrate that monoperphthalic acid is consumed by reaction with the ethylenic linkage, Chakravorty and Levin 25 were the first to isolate oxiranes by the oxidation of unsaturated compounds with this oxidizing agent. Epoxidation with monoperphthalic acid is conducted under the same conditions as with perbenzoic acid, and good yields of oxiranes are obtained. Epoxidation with monoperphthalic acid has been applied most extensively to naturally occurring products, such as sterols and polyenes. Ethylenic compounds which have been converted to oxiranes by epoxidation with monoperphthalic acid are listed in Table I.

Peracetic Acid. Since peracetic acid is one of the most conveniently prepared organic peracids, a study of its possible use as an epoxidizing agent was to be expected. For a long time, however, it was assumed that oxiranes could not be prepared by the epoxidation of olefins with peracetic acid since the products isolated from such reactions were either  $\alpha$ -glycols or their monoacetates. The first successful epoxidation with peracetic acid was reported by Böeseken, Smit, and Gaster, 28,29 who obtained methyl 9,10,12,13-diepoxystearate from methyl linoleate, but the yields were extremely poor and the major proportion of the product consisted of a polymer of undetermined constitution.³⁰ In a systematic study of the reaction of unsaturated compounds with peracetic acid in

²³ Findley and Swern, U. S. pat. 2,567,930 [C. A., 46, 3560 (1952)].

²⁴ Baeyer and Villiger, Ber., 34, 762 (1901).

²⁵ Chakravorty and Levin, J. Am. Chem. Soc., 64, 2317 (1942).

²⁶ Böhme, Ber., 70, 379 (1937).

²⁷ Böhme and Steinke, Ber., 70, 1709 (1937). ²³ Böeseken, Smit, and Gaster, Proc. Acad. Sci. Amsterdam, 32, 377 (1929).

²⁹ Smit, Rec. trav. chim., 49, 675 (1930).

³⁰ Swern, unpublished results.

acetic acid solution and in inert solvents, Arbusow and Michailow 31,32 observed that hydroxy acetates were formed in acetic acid while good yields of oxiranes were obtained in inert solvents. They concluded that the behavior of peracetic acid toward olefins is the same as that of perbenzoic acid, but that when an acetic acid solution is employed the oxirane is converted to the hydroxy acetate by further reaction with acetic acid. The apparent necessity for employing peracetic acid in an inert solvent to obtain good yields of oxiranes discouraged the general use of peracetic acid for epoxidation, because peracetic acid can be prepared and used most conveniently in acetic acid, whereas its isolation free (or substantially free) of acetic acid is time-consuming and hazardous.

Subsequently, however, in connection with a kinetic study of the reaction of peracetic acid in acetic acid solution with various long-chain olefins, suitable reaction conditions were determined for the efficient conversion of ethylenic compounds to oxiranes.1 To obtain good yields of oxiranes it is necessary to operate at moderate temperatures (20-25° is preferred), to keep the reaction time as short as possible and to exclude strong acids, which catalyze the opening of the oxirane ring by acetic acid. The reaction was shown to be general and afforded a simple and convenient method for the preparation of oxirane compounds in quantity. Isolation of pure peracetic acid and employment of inert solvents were unnecessary. Yields of oxiranes, however, were usually lower than when perbenzoic or monoperphthalic acid was employed. In the peracetic acid epoxidation of compounds containing both an ethylenic and an acetylenic linkage, it has been reported that only the double bond is attacked.33,34 Acetylenic compounds react with peracetic acid, but the rates of reaction are only about one-thousandth as great as the rates of reaction of analogous ethylenic compounds. Three atoms of oxygen add, and the acetylenic linkage is cleaved. Oxirenes, -C----C-, are

intermediates and have been isolated from some reactions.34a

Ethylenic compounds which have been converted to oxiranes by epoxidation with peracetic acid are listed in Table I.

Percamphoric Acid. Percamphoric acid has been employed to convert ninene and cholesterol to the corresponding oxiranes.35

³¹ Arbusow and Michailow, J. prakt. Chem., 127, 1 (1930). 22 Arbusow and Michailow, J. prakt. Chem., 127, 92 (1930).

³³ Malenok and Sologub, J. Gen. Chem. (U.S.S.R.), 10, 150 (1940) [C. A., 34, 7286

Malenok and Sologub, J. Gen. Chem. (U.S.S.R.), 11, 983 (1941) [C. A., 37, 355 (1943)].

³⁴a Schlubach and Franzen, Ann., 577, 60 (1952).

Milas and Cliff, J. Am. Chem. Soc., 55, 352 (1933).

Performic Acid. Performic acid is generally considered not to be an epoxidation reagent because the high acidity of formic acid (employed either as solvent or formed in the oxidation) causes most oxirane rings to open rapidly. It has been shown recently, however, that  $\alpha$ -diisobutylene yields an isolable oxirane on oxidation with performic acid, although the yield is low.36 By employing only small quantities of formic acid as solvent and oxygen carrier, and in some cases by adding small amounts of sodium hydroxide, it has been reported that methyl oleate, octyl oleate, propylene glycol dioleate, and soybean oil can be converted to oxiranes in fair yields.²⁷ Recently, two steroids have been converted to oxiranes by epoxidation with performic acid.38,39

The diisobutylenes behave somewhat abnormally on reaction with both performic and peracetic acids, yielding, besides the expected products, unsaturated alcohols, an aldehyde, a ketone, a cyclic diether, and high-boiling products. 36, 40-43

#### Hydroxylation

Peracetic Acid. The use of peracetic acid for the preparation of  $\alpha$ -glycols from unsaturated substances probably exceeds that of all other organic peracids combined. Peracetic acid is usually prepared and employed in either of two ways: (1) the peracid is preformed by the reaction of acetic acid or acetic anhydride with 25-90% hydrogen peroxide 1,44-47 and then mixed with the unsaturated compound, or (2) the unsaturated compound is mixed with hydrogen peroxide and acetic acid, and the peracetic acid is consumed as it is formed.4,48 Under suitable conditions (p. 381) oxiranes are obtained in good yields; but in the manner that the reactions have usually been carried out (long reaction times, and/or high temperatures, and/or in the presence of sulfuric acid), the products isolated are hydroxy acetates formed by the reaction of excess acetic acid with the oxirane produced initially. The hydroxy

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<sup>26</sup> Byers and Hiekinbottom, J. Chem. Soc., 1948, 1328.
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³⁷ Niederhauser and Koroly, U. S. pat. 2,485,160 [C. A., 44, 7346 (1950)].

³⁸ Djerassi, Mancera, Stork, and Rosenkranz, J. Am. Chem. Soc., 73, 4496 (1951). ³⁹ Stork, Romo, Rosenkranz, and Djerassi, J. Am. Chem. Soc., 73, 3546 (1951).

⁴⁰ Byers and Hiekinbottom, J. Chem. Soc., 1948, 284.

⁴¹ Byers and Hickinbottom, Nature, 158, 341 (1946).

⁴² Hickinbottom, J. Chem. Soc., 1948, 1331.

⁴³ Hiekinbottom, Nature, 159, 844 (1947).

⁴⁴ D'Ans and Frey, Ber., 45, 1845 (1912). 45 D'Ans and Frey, Z. anorg. Chem., 84, 145 (1914).

⁴⁵ D'Ans and Kneip, Ber., 48, 1136 (1915). Greenspan, J. Am. Chem. Soc., 68, 907 (1946).

⁴ Greenspan, Ind. Eng. Chem., 39, 847 (1947).

acctates are readily hydrolyzable to  $\alpha$ -glyeols in excellent yield.⁴⁹⁻⁵² Although good yields of glycols were reported by some early investigators, the operating conditions employed eaused the loss of much active oxygen by decomposition. With sulfurie acid as the catalyst, moderate temperatures (40°), and short reaction periods, excellent yields of  $\alpha$ -glycols are obtained with stoichiometric quantities of 25–30% hydrogen peroxide.⁴ Since the sulfuric acid catalyzes the formation of peracetic acid and the peracid is rapidly consumed at 40°, the reaction is complete in a few hours and little active oxygen is lost. This procedure is one of the most efficient for converting long-chain olefins to  $\alpha$ -glycols. Slightly higher yields of  $\alpha$ -glycols are obtained when 90% hydrogen peroxide is employed.⁴⁸

Ethylenic compounds which have been converted to  $\alpha$ -glycols by oxidation with peracetic acid, either preformed or prepared and utilized in situ, are listed in Table I. Some of the unsaturated substances listed have been converted to hydroxy acetates rather than to  $\alpha$ -glycols, but the conversion to glycols is effected so readily by hydrolysis that these substances have also been included.

Performic Acid. An even more efficient and rapid hydroxylation technique eonsists in the reaction of unsaturated compounds with performic acid. Not only is performic acid formed rapidly when 25-90% hydrogen peroxide and formic acid are mixed,44-47,53 but it also reacts rapidly and completely with the unsaturated linkage. By means of this hydroxylation reaction, conversion of an unsaturated compound to an α-glyeol is accomplished within a short time, and approximately stoichiometric quantities of hydrogen peroxide can be employed. The initial product of oxidation is not the  $\alpha$ -glycol but the oxirane, which is rapidly converted in most eases to a hydroxy formate as a result of the high acidity of formie acid. Hydroxy formates are the products usually isolated and are readily converted to the a-glyeols by hydrolysis with dilute aqueous alkali or even by exposure to moist air or heating with water.1 It is important to note that performie acid is preferably not prepared separately, because it is unstable and loses oxygen rapidly, 46,47,63,64 but it is prepared and utilized in situ.4 Somewhat more complete hydroxylation is obtained by employing 90% hydrogen peroxide instead of the 25-30% concentration.48

Concentrated solutions of performic acid can be used in the hydroxyl-

et Hildsteh, J. Chem. Soc., 1926, 1828.

¹ Hilditch and Lea, J. Chem. Soc., 1927, 3106.

⁴¹ Scanlan and Swern, J. Am. Chem. Soc., 62, 2305 (1940).

 ¹³ Scanlar, and Swern, J. Am. Chem. Soc., 62, 2309 (1940).
 ¹⁴ Tomnies and Homiller, J. Am. Chem. Soc., 64, 3054 (1942).

[#] Swern and Findley, unpublished results.

ation of  $\alpha,\beta$ -unsaturated acids to give fair yields of dihydroxy acids within a relatively short time. 55 Dilute solutions of organic peracids either are ineffective in hydroxylation of such compounds, or extremely long reaction times are required during which loss of active oxygen occurs.

The performic acid oxidation of ethylenic compounds having a hydroxyl group on a carbon atom directly adjacent to the ethylenic group yields appreciable amounts of acidic chain cleavage products in addition to about 50% of the expected hydroxylation products.56

In the peracetic and performic acid hydroxylation of compounds containing both an ethylenic and an acetylenic linkage only the double bond is attacked.34a, 57-60

Ethylenic compounds converted to  $\alpha$ -glycols by oxidation with performic acid are listed in Table I.

Perbenzoic, Monoperphthalic, or Percamphoric Acid. These acids can be employed for the preparation of  $\alpha$ -glycols from olefins by hydrolyzing the oxiranes which are formed first. In general, there is no advantage in employing the aromatic peracids to prepare  $\alpha$ -glycols when two more-efficient peracids (performic and peracetic acid) are available for this purpose. In the presence of water or with unusually long reaction times, reactions have been reported in which  $\alpha$ -glycols or their monobenzoates rather than oxiranes were obtained from oxidations of olefins with perbenzoic acid.

Ethylenic compounds which have been converted to  $\alpha$ -glycols or to hydroxybenzoates by oxidation with perbenzoic acid are listed in Table I.

#### STEREOCHEMISTRY AND MECHANISM

Although the structure of organic peracids, usually written RCO₃H, is not known, it is evident from their numerous and varied reactions that they are electrophilic reagents. 10 As the nucleophilic nature of an olefin is increased by replacement of the hydrogen atoms of its ethylenic linkage with electron-releasing groups, the rate of reaction with organic peracids increases considerably (see p. 388). Since peracid reactions investigated so far are subject to general acid catalysis, 61,62 it has been

⁵⁵ English and Gregory, J. Am. Chem. Soc., 69, 2120 (1947).

⁵⁶ Ross, Gebhart, and Gerecht, J. Am. Chem. Soc., 71, 282 (1949).

Evans, Fraser, and Owen, J. Chem. Soc., 1949, 248. ⁵⁸ Malenok, J. Gen. Chem. (U.S.S.R.), 9, 1947 (1939) [C. A., 34, 4385 (1940)].

маненок, J. Gen. Chem. (О.В.Б.), 5, 1904 (1936) [C. A., 31, 4285]

⁵⁹ Malenok and Sologub, J. Gen. Chem. (U.S.S.R.), 6, 1904 (1936) [C. A., 31, 4285]

⁶⁰ Raphael, J. Chem. Soc., 1949, S44.

⁶¹ Friess, J. Am. Chem. Soc., 71, 2571 (1949).

⁶² Waters, J. Chem. Soc., 1948, 1574.

proposed that the attacking moiety in peracid oxidations is the electropositively polarized (electrophilic) hydroxyl group [O:H]+.63,64 reaction of an olefin, such as propylene, with a peracid may, therefore, be represented as follows.10

$$\begin{array}{c} CH_3 \rightarrow -CH = \overset{\frown}{C}H_2 + \overset{\frown}{[0:H]^+} \rightarrow \\ [CH_3 \rightarrow -CH - CH_2] \rightarrow CH_3 - CH - CH_2 + H^+ \\ \vdots \vdots \vdots \\ H \end{array}$$

This simple formulation, however, does not account for the striking stereospecificity of the reaction which precludes a free carbonium ion intermediate. A more reasonable alternative mechanism would involve essentially direct formation of the conjugate acid of the oxirane by donation of [O:H]+ to the olefin by a peracid-general acid complex in a manner similar to that shown in the accompanying equation. The olefin-

$$\begin{array}{c|c} H_3C \\ HC & H & O \\ \parallel & O - O - C - R \end{array} \rightarrow \begin{bmatrix} CH_3 \\ \mid & CH \\ CH \end{bmatrix}^+ + \begin{bmatrix} O \\ \parallel & CH \\ R - C - O - HA \end{bmatrix}^- \rightarrow \\ CH_3 - CH - CH_2 + RCO_2H + HA \end{array}$$

[O:H]⁺ part of the transition state of such a process would be similar to the so-called  $\pi$ -complexes. This mechanism obviates any necessity for postulation of rapid and reversible [O:H]+ formation from peracid and general acid (HA) followed by a slow attack of [O:H]+ on the double bond. It is also a more reasonable reaction path in the non-polar solvents often used as reaction media.

As discussed earlier (pp. 380-385) the product isolated may be the

ti Weisenborn and Taub, J. Am. Chem. Soc., 74, 1329 (1952). a Roitt and Waters, J. Chem. Soc., 1949, 3060.

M. J. S. Dewar, The Electronic Theory of Organic Chemistry, Oxford University Press, 1949.

oxirane or the hydroxy acyloxy compound, depending on the experimental conditions, the peracid used, and the stability of the oxirane.

The initial oxidation step in epoxidation and hydroxylation with organic peracids is the same, and it has generally been assumed that this reaction proceeds by *cis* addition to the double bond.^{6,66} Recently, unequivocal evidence was obtained to substantiate this assumption. It was shown by x-ray diffraction and infrared absorption studies that oleic acid and oleyl alcohol (both *cis* olefins) yield *cis*-9,10-epoxystearic acid and *cis*-9,10-epoxyoctadecanol, respectively, on epoxidation with peracetic or perbenzoic acid, and the corresponding *trans* olefins, elaidic acid and elaidyl alcohol, yield *trans*-9,10-epoxystearic acid and *trans*-9,10-epoxyoctadecanol, respectively.⁶⁷

Opening of the oxirane ring, in the preparation of  $\alpha$ -glycols from the corresponding oxiranes, is accompanied by inversion whether the reaction is conducted in neutral, acidic, or alkaline media.⁵ The only exception to this generalization apparently is the opening of an oxirane ring in the terminal position of an aliphatic chain. In this case, if the ring-opening reagent attacks the terminal position, inversion cannot occur. A reaction scheme correlating the configurational relationships in the conversion of oleic and elaidic acids (cis- and trans-9-octadecenoic acids, respectively) to 9,10-dihydroxystearic acids by way of the intermediate oxiranes has recently been published.⁵ This scheme is self-consistent and is in harmony with accepted theories of inversions, double-bond addition reactions, and the vast amount of experimental data available. This reaction sequence is undoubtedly of general applicability to other olefins with non-terminal double bonds.

It should be noted that the oxirane obtained by epoxidation of an olefin with organic peracids (cis addition) is identical with that obtained by with organic peracids (cis addition) followed by dehydrohalogenation (inhypohalogenation (trans addition) followed by dehydrohalogenation (inhypohalogenation (trans addition) followed by dehydrohalogenation (inhypohalogenation). In the latter preparative procedure two inversions have version occurs). In the latter preparative procedure two inversions.

Hydroxylation of olefins with potassium permanganate, 70-73 t-butyl hydroperoxide (osmium tetroxide catalyst), 74, 75, 76 or by photochemical

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Braun, J. Am. Chem. Soc., 51, 228 (1929).
Witnauer and Swern, J. Am. Chem. Soc., 72, 3364 (1950).
Abderhalden and Eichwald, Ber., 48, 1847 (1915).
Sowden and Fischer, J. Am. Chem. Soc., 64, 1291 (1942).
Böeseken, Rec. trav. chim., 47, 683 (1928).
Böeseken and Cohen, Rec. trav. chim., 47, 839 (1928).
King, J. Chem. Soc., 1943, 37.
Kuhn and Ebel, Ber., 58, 919 (1925).
Kuhn and Ebel, Ber., 58, 919 (1925).
Milas, J. Am. Chem. Soc., 59, 2342 (1937).
Milas and Sussman, J. Am. Chem. Soc., 58, 1302 (1936).
Milas, Sussman, and Mason, J. Am. Chem. Soc., 61, 1844 (1939).
Milas, Sussman, and Mason, J. Am. Chem. Soc., 61, 1844 (1939).
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addition of hydrogen peroxide to the double bond ⁷⁷ proceeds by *cis* addition. Catalytic hydroxylation of olefins with hydrogen peroxide and other inorganic catalysts, such as pertungstic acid, pervanadic acid, or selenium dioxide, however, proceeds by *trans* addition.⁷⁸

#### SELECTION OF EXPERIMENTAL CONDITIONS

Since the oxirane group is extremely reactive and undergoes ring opening with various types of compounds which contain active hydrogen atoms, it is obvious that conditions for epoxidation must be selected with care. It is of paramount importance to avoid high reaction temperatures ¹ and to exclude strongly acidic materials from the reaction mixtures ⁴ if high yields are to be obtained. In epoxidations with perbenzoic and monoperphthalic acids an inert solvent is employed; in epoxidations with peracetic acid, acetic acid may be used as the solvent, provided that strong acids are absent and reaction temperatures below about 30° are employed.

With unsaturated substances containing isolated double bonds, such as 2-pentene, 2-butene, oleic acid, and oleyl alcohol, epoxidation is rapid and is usually complete within eight to twenty-four hours at room temperature or below. If electron-releasing groups are attached to or are in close proximity to the ethylenic linkage, as in 2-methylpropene, 2methyl-2-butene, and tetramethylethylene, the reaction is considerably accelerated; 10 if electron-attracting groups are attached to or are in close proximity to the ethylenic linkage, as in cinnamic, maleic, fumaric, crotonic, 2-pentenoic, and 2-hexenoic acids and their esters, the reaction is slowed down.10 The wide range of specific reaction rates in related groups of compounds is shown most strikingly by comparing ethylene  $(k \times 10^3 = 0.19)$  with 2-methyl-2-butene  $(k \times 10^3 = \text{ca. } 1000)$ , cyclobutene  $(k \times 10^3 = 21)$  with 1-methylcyclopentene  $(k \times 10^3 = 2200)$ , sorbic acid ( $k \times 10^3 = 0.04$ ) with oleic acid ( $k \times 10^3 = 384$ ), allylbenzene  $(k \times 10^3 = 2.0)$  with 1-phenyl-1-propene  $(k \times 10^3 = 46)$ , 1,4-dihydronaphthalene ( $k \times 10^3 = 37$ ) with 1,2-dihydronaphthalene  $(k \times 10^3 = 230-240)$ , cinnamic acid  $(k \times 10^3 = 0.13)$  with cinnamyl alcohol (k  $\times$  10³ = 203), 1-phenyl-2-butene (k  $\times$  10³ = 10) with 1phenyl-1-butene ( $k \times 10^3 = 80$ ), cugenol ( $k \times 10^3 = 2.2$ ) with isoeugenol ( $k \times 10^3 = 127$ ), and safrole ( $k \times 10^3 = 1.3$ ) with isosafrole  $(k \times 10^3 = 148).^{10.79}$  Furthermore, the specific reaction rate of tetramethylethylene with peracetic acid at 25.8° is too high to be meas-

¹⁷ Milas, Kurz, and Anslow, J. Am. Chem. Soc., 59, 543 (1937).

⁷¹ Mugdan and Young, J. Chem. Soc., 1949, 2988. 71 Swern, Chem. Revs., 45, 1 (1949).

Selected references describing kinetic studies are 2, 28, and ured.80,81 80-88.

The rates of oxidations with peracids can be determined readily with a minimum of experimental effort by measuring unconsumed peroxide at suitable time intervals.11,13,89,90,91 By following the disappearance of active oxygen, the reaction can be terminated at exactly the right time, thereby minimizing side reactions and loss of active oxygen. Furthermore, the determination of unconsumed peroxide should be carried out in all peracid oxidations in which distillation techniques are employed in the recovery of solvent and in the isolation of reaction products. In reactions which proceed slowly, a large amount of unconsumed peracid may be present in the distillation charge and cause an explosion if the peroxide is not destroyed.

Although a wide range of conditions can be employed in the preparation of  $\alpha$ -glycols, temperatures above 50° are undesirable because significant loss of active oxygen occurs. Early workers, who were not concerned with efficient use of active oxygen, operated at high temperatures and of necessity employed large excesses of hydrogen peroxide or peracid. Reaction temperatures below 5-10° may also be disadvantageous since they make the reaction time objectionably long.

To help in the sclection of hydroxylation techniques, the methods just discussed are listed in decreasing order of efficiency and over-all desirability from the laboratory standpoint.

- 1. Oxidation with 30% hydrogen peroxide in formic acid solution at 40°; 1.025-1.05 moles of hydrogen peroxide per ethylenic linkage.2,4 This method is admirably suited for the hydroxylation of isolated double bonds and is probably the best hydroxylation technique employing organic peracids. See also method 3.
  - 2. Oxidation with 30% hydrogen peroxide in acetic acid solution containing catalytic quantities of sulfuric acid at 40°; 1.025-1.05 moles of hydrogen peroxide per ethylenic linkage.2.4

⁸⁰ Böeseken and Stuurman, Proc. Acad. Sci. Amsterdam, 39, 2 (1936) [C. A., 30, 3304] (1936)1.

⁸¹ Böeseken and Stuurman, Rec. trar. chim., 56, 1034 (1937).

⁸² Bodendorf, Arch. Pharm., 268, 491 (1930).

⁸³ Böeseken and Blumberger, Rec. trav. chim., 44, 90 (1925).

⁸⁴ Böeseken and Hanegraaff, Rec. trav. chim., 61, 69 (1942). Boeseken and Hanegraan, Acad. Sci. Fennicae, A59, No. 13, 3 (1943) [C. A., 41, 2307 (1947)]. B Heinänen, Ann. Acad. Sci. Fennicae, A59, No. 13, 3 (1943) [C. A., 41, 2307 (1947)].

Sinit, Rec. trav. chim., S. Amsterdam, 38, 450 (1935) [C. A., 29, 4657 (1935)]. Stuurman, Proc. Acad. Sci. Amsterdam, 1996.

^{5.} Stuurman, tnesis, Oliversis, Vol. II, 2nd ed., p. 413, Springer, Berlin, 1931.
55 Kolthoff and Menzel, Die Massanalyse, Vol. II, 2nd ed., p. 413, Springer, Berlin, 1931. 8 J. Stuurman, thesis, University of Delft, 1936.

³⁰ Marks and Morrell, Analyst, 54, 503 (1929).

⁹¹ Wheeler, Oil and Soap, 9, 89 (1932).

- 3. The same as 1 and 2, but employing 90% hydrogen peroxide. 48, 55 Although slightly more complete reaction is obtained with 90% hydrogen peroxide, the hazards attendant upon its use make it less desirable for laboratory investigation. 92, 93, 94 By use of the more concentrated peracids, however, ethylenic linkages adjacent to earboxyl groups can be hydroxylated readily.55
- 4. Prior preparation of performic or peracetic acids and employment of the peracids under conditions similar to 1, 2, and 3 above.
- 5. Epoxidation with peracetic, perbenzoic, or monoperphthalic acid, followed by hydrolysis. The only virtue of this technique, probably, is that one can obtain either the oxirane or the  $\alpha$ -glycol from a given unsaturated substance.

Because of the instability of performic acid, there is usually little point in its separate preparation (method 4). If it is prepared separately, however, it should be used immediately. Performic acid of 90% strength is highly explosive. 94a In contrast to performic acid, peracetic acid is relatively stable and can be stored conveniently. In the absence of eatalysts, concentrated solutions of peracetic acid are fairly stable at room temperature (15-25°); 87-95% solutions remain virtually unchanged on standing for about five weeks,46 the 50% solution shows no loss of peracid after storage for two weeks, 45 and the 45% solution retains 75% of the peracid after seven weeks. The 45% solution retains 94% of the peracid after seven weeks of storage if it is stabilized with sodium pyrophosphate 47 (other stabilizers have also been suggested). 95,96 Five to ten per cent solutions of peracetic acid in acetic acid, however, show significant losses of active oxygen at room temperature but little loss at 0 to 5°.1 Although peracetic acid can be prepared by efficient processes and only a small amount of active oxygen is lost or unavailable for oxidative purposes, the separate preparation of the peracid is a time-consuming step in the hydroxylation reaction, and method 2 is more desirable. Concentrated solutions of peracetic acid have recently become commercially available.97

There is a wide variety of methods for preparing organic peracids, and many solvents have been suggested for use in their preparation, isolation, and application as oxidizing agents. This phase of peracid chemistry is

[&]quot; Bellinger, Friedman, Bauer, Eastes, and Bull, Ind. Eng. Chem., 38, 310 (1946).

Bellinger, Friedman, Bauer, Eastes, and Edmonds, Ind. Eng. Chem., 38, 627 (1946). ²⁴ Shanley and Greenspan, Ind. Eng. Chem., 39, 1536 (1947).

we Weingartshofer-Olmos and Gigubre, Chem. Eng. News, 30, 3041 (1952).

n Naamloore Venootschap Industrieele Maatschappij Voorheen Noury and Van Der Lan le and Van Der Lande, Brit, pat. 234,163 [C. A., 20, 768 (1926)].

r Reichert, McNeight, and Elston, U. S. pat. 2,347,434 [C. A., 39, 89 (1945)]. Buffalo Electrochemical Co., Peracetic Acid Data Sheet 1 (1947).

not sufficiently pertinent to be discussed here in detail, but information has recently been published on this subject.79 The particular oxidative method and solvent selected will depend, in large part, on the solubility of the peracid and on the structure of the unsaturated substance and the oxidation products. Furthermore, the stability of the peracid and the oxidation products in the solvent medium and the ease of separation of the desired products from the other materials present have an important bearing on the selection of reaction conditions. The solvent has been reported to affect the rates of decomposition of peracids as well as their rates of reaction with unsaturated substances.7,13,83,98-101

For information regarding other organic peracids (properties, methods of preparation, special techniques, etc.) reference 79 can be consulted.

#### EXPERIMENTAL PROCEDURES

Caution. All preparations of and reactions with organic peracids should be conducted behind a safety shield, because a reaction occasionally proceeds with uncontrollable violence. When an olefin of unknown structure or one that contains at least three electron-releasing groups attached to or in close proximity to the ethylenic linkage is epoxidized or hydroxylated for the first time, the reaction should be run on a small scale (preferably 0.1 mole or less), and provision should be made for efficient cooling. Detailed information regarding the properties of concentrated hydrogen peroxide 92, 93, 94, 102-105 and organic peracids 79 has recently been published.

Peracid oxidation mixtures should not be distilled unless an analysis has indicated the absence or low concentration of active oxygen. When the peracid content is low, acetic and formic acids can be safely and completely distilled from oxidation reactions at or below room temperature by the use of low pressure. Peracids and other peroxides can be conveniently destroyed by the addition of ferrous sulfate, sodium bisulfite, or other reducing agents.

⁹⁹ Berezovskaya and Semikhatova, Bull. acad. sci. U.R.S.S., Classe sci. math. nat., 1934, 1583, 1589 [C. A., 29, 6130 (1935)]. ³⁹ Calderwood and Lane, J. Phys. Chem., 45, 108 (1941).

¹⁰⁰ Lagrave, Ann. Chim., [10] 8, 363 (1927).

¹⁰¹ Meerwein, Ogait, Prang, and Serini, J. prakt. Chem., 113, 9 (1926). 102 Bretschger and Shanley, Trans. Electrochem. Soc., 92, 10 pp. (1947) preprint.

¹⁰³ McKee, Mech. Eng., 68, 1045 (1946).

¹⁰⁴ Médard, Compt. rend., 222, 1491 (1946).

¹⁰⁵ Schumb, Ind. Eng. Chem., 41, 992 (1949).

#### Analysis of Peracids

Perbenzoie Aeid. Perbenzoie acid in an organie solvent can be determined iodimetrically by shaking the solution with an aqueous acetie acid solution of potassium iodide. A known volume of the perbenzoic acid solution is pipetted into an iodine flask containing 50 ml. of 0.4 N acetie acid and 1 g. of potassium iodide, the mixture is shaken, and the liberated iodine is titrated with 0.05-0.1 N sodium thiosulfate solution, starch indicator being used.

In following the course of the oxidation of water-insoluble substances which precipitate upon addition of the solution to the aqueous acetie acid, a sharper end point is obtained by adding the perbenzoic acid solution to 25 ml. of a chloroform-acetic acid solution (3:2 by volume). Two milliliters of saturated potassium iodide solution is added, and the mixture is allowed to stand for five minutes. Seventy-five milliliters of water is added, the solution is shaken, and the liberated iodine is titrated with 0.05-0.1 N sodium thiosulfate.91 One milliliter of 0.1 N sodium thiosulfate is equivalent to 0.00690 g. of perbenzoic acid.

Monoperphthalic Acid. Monoperphthalic acid can be determined by the same methods employed for perbenzoic acid. An alternative procedure 106 is to add 2 ml. of the solution to 30 ml. of 20% aqueous potassium iodide and titrate the liberated iodine after 10 minutes with 0.05 N sodium thiosulfate solution. One milliliter of 0.05 N sodium thiosulfate is equivalent to 0.00455 g. of monoperphthalic acid.

Peracetic Acid. The peroxide components in the peracetic acid solutions described below are determined on a single sample as follows: 44,45 0.2-2 ml. of the solution (accurately dispensed from a pipette or weighed) is diluted with 50 ml. of 4 N aqueous sulfurie acid which has been cooled to 0°. This solution is titrated rapidly with  $0.1\,N$  potassium permanganate to a pink end point. This determines unreacted hydrogen peroxide; 1 ml. of 0.1 N potassium permanganate is equivalent to 0.00170 g. of hydrogen peroxide. The peracetic acid is determined by adding 2 ml. of saturated aqueous potassium iodide to the same solution and rapidly titrating with 0.1 N sodium thiosulfate, starch indicator being used; 1 ml. of 0.1 N sodium thiosulfate is equivalent to 0.00380 g. of perneetic acid. At this point, the flask and its contents are heated on the steam bath for five to ten minutes, eausing a return of the blue color, and liberated iodine is titrated with 0.1 N sodium thiosulfate. The last titration gives the diaeetyl peroxide content; 1 ml. of 0.1 Nsodium thiosulfate is equivalent to 0.00590 g. of diaeetyl peroxide.

¹²⁴ B5hme, Org. Syntheses, 20, 70 (1940).

has been reported that ceric sulfate is more satisfactory than potassium permanganate for determination of residual hydrogen peroxide. 107

In following the consumption of active oxygen during the oxidation of water-insoluble substances with peracetic acid, the procedure described under the analysis of perbenzoic acid should be employed.91 This determines total active oxygen and not peracetic acid alone, but the difference between the titrations at succeeding time intervals gives a measure of peracetic acid consumed.

Performic Acid. The procedures described in the analysis of peracetic acid are used.

#### Preparation of Peracids

Perbenzoic Acid (Benzoyl Peroxide-Sodium Methoxide Method). Directions published in Organic Syntheses 11 are probably the most satisfactory for preparing stable solutions of perbenzoic acid. Briefly, this method consists in (a) allowing benzoyl peroxide to react with sodium methoxide in chloroform-methanol solution, (b) extracting the sodium perbenzoate solution with water, (c) acidifying with sulfuric acid, and (d) extracting the perbenzoic acid with chloroform. Yields of perbenzoic acid of about 85% are obtained. Do not recrystallize benzoyl peroxide from hot chloroform, as suggested in the original Organic Syntheses procedure, as this operation is hazardous. Benzoyl peroxide may be purified safely by adding methanol to a chloroform solution of the peroxide at room temperature. 108 A recrystallized grade is commercially

For preparation of large quantities of perbenzoic acid or solutions which are to be stored for a long time, a modified procedure has been

- (a) The mixture is kept below 0° during the addition of the chloroform solution of benzoyl peroxide to the methanol solution of sodium recommended.13 methoxide. Since this reaction is highly exothermic, a large quantity of salt-ice freezing mixture at  $-15^{\circ}$  is employed to cool the reaction flask, the benzoyl peroxide solution is added at a slow, even rate of about 15-20 ml. per minute, and the reaction flask is swirled vigorously and continuously during the addition. There is no need to wait four to five minutes, as specified in the original procedure 11 before extracting the mixture with water.
  - (b) Instead of transferring the chloroform-methanol solution containing sodium perbenzoate to a separatory funnel, about 150 ml. of

¹⁰⁷ Greenspan and MacKellar, Anal. Chem., 20, 1061 (1948). 108 Nozaki and Bartlett, J. Am. Chem. Soc., 68, 1686 (1946).

¹⁰⁹ Lucidol Corporation, Buffalo, New York.

water containing chopped ice is added to the reaction mixture which is rapidly swirled. The mixture is then transferred to the separatory funnel, and 350 ml. of water containing chopped ice is added to the rapidly swirled material. In this way, the formation of lumps which dissolve slowly is prevented.

(c) The emulsion that collects at the interface of the aqueous sodium perbenzoate phase and the chloroform phase is discarded. Only three to five minutes is allowed for separation of the phases. Likewise, emulsions formed during the washing of the aqueous layer are discarded.

(d) The aqueous phase is washed with two 100-ml. portions of carbon

tetrachloride, instead of chloroform.

(e) After acidification, the aqueous solution is extracted with reagent-grade benzene rather than chloroform. At this point, the temperature of the solution should be above 5°, to prevent freezing of the benzene.

(f) The benzene solution is washed with water, dried over anhydrous sodium sulfate (calcium chloride sometimes causes a sudden decomposition of the peracid 11), and stored in the dark at about 10° until used.

Crystalline perbenzoic acid can be obtained by removal of the solvent under vacuum, as described in *Organic Syntheses*, ¹¹ and purified by recrystallization from chloroform-ethanol mixtures ¹¹⁰ or from petroleum ether. ¹¹¹ Perbenzoic acid melts at about 41° and is soluble in the common organic solvents, except cold petroleum ether.

Perbenzoic Acid (Benzaldehyde-Air Method).³ The air oxidation of benzaldehyde in acetone solution irradiated with ultraviolet light is a convenient method for the preparation of moderately large quantities of perbenzoic acid.

In a 5-l. three-necked Pyrex flask equipped with a thermometer, a solid carbon dioxide-cooled reflux condenser, and two fritted glass disks reaching to the bottom of the flask, 520 g. (4.9 moles) of freshly distilled benzaldehyde is dissolved in 4 l. of acetone. The flask is immersed in an ice-water bath and irradiated from the top with three 125-watt Hanovia quartz mercury-vapor lamps, symmetrically placed around the flask, while a rapid stream of dry air is passed through the fritted disks and into the solution for twenty-four hours at 5-10°. The reaction is conducted in a fume hood because of the formation of ozone. If the reaction cannot be run without interruption, the acetone solution can be stored at 5-10° with little or no loss of perbenzoic acid. After about twenty-four hours, the rate of peracid formation decreases considerably

¹¹⁰ Maan, Rec. trav. chim., 48, 332 (1929).

¹¹¹ Baeyer and Villiger, Ber., 33, 1569 (1900).

and the solution then contains about 2 moles of perbenzoic acid. The yield is 40-45%.

Monoperphthalic Acid. The procedure described in Organic Syntheses,  106  consisting in the reaction of phthalic anhydride with alkaline 30%aqueous hydrogen peroxide, is satisfactory, and gives 65-70% yields. It has been reported to be advantageous to employ 40% sodium hydroxide solution and to add crushed ice directly to the reaction mixture. 112 In this procedure, the peracid is extracted with ether, but, if ether is not a suitable solvent for the subsequent oxidation reactions, it can be removed readily and replaced by dioxane or other solvent by a procedure described in Organic Syntheses. 106

Peracetic Acid. 1,47 In a 5-l. three-necked flask equipped with a mechanically driven glass stirrer, a thermometer, and a separatory funnel is placed 2250 g. of acetic anhydride, which has been filtered through glass wool to remove particles which may catalyze peroxide decompo-The thermometer should be immersed in the liquid, and at least one neck of the flask should be open to the atmosphere. The acetic anhydride is warmed to 35-40° in a water bath into which cold or warm water can be run at will and removed rapidly if necessary. By means of the separatory funnel, 500 g. of 25-30% hydrogen peroxide is added in about one hour with agitation, the temperature being maintained at 40°. The reaction becomes mildly exothermic soon after the addition of hydrogen peroxide is started, and cooling is required for three to four hours after the addition is complete to maintain the temperature at 40° (bath temperature 25-30°). The solution is allowed to stand overnight at room temperature. The concentration of peracetic acid is then about 0.8–1.2 M (6–9%). The yield is 60–90%. solution contains diacetyl peroxide and some unconverted hydrogen peroxide in addition to peracetic acid and acetic acid.

A concentrated solution of peracetic acid 47 is prepared by cautiously adding 9.1 g. of 90% hydrogen peroxide to a stirred solution of 10 g. of acetic acid and 0.11 ml. of concentrated sulfuric acid contained in a flask immersed in a water bath at 22-23°. At the end of four hours, the Peracetic acid content of the solution is about 44%; it rises to a maximum of 46% within twelve to fifteen hours.

Performic Acid. 47,53,54 In a 500-ml. Erlenmeyer flask, 25 g. of 25-30% hydrogen peroxide and 250 g. of 98-100% formic acid are mixed at room temperature. Since the reaction is only mildly exothermic (temperature rise 1-2°), no cooling is required in batches of this size. The maximum content of performic acid (approximately 5%) is obtained within thirty

¹¹² Bachman and Cooper, J. Org. Chem., 9, 302 (1944).

to sixty minutes, as determined by the analytical techniques already described.

A concentrated solution of performic acid is prepared by cautiously adding 28.4 g. of 90% hydrogen peroxide to a stirred solution of 23.0 g. of 98–100% formic acid and 0.28 ml. of concentrated sulfuric acid contained in a flask immersed in a water bath at 22–23°.47,55 Maximum performic acid concentration (approximately 35%) is reached within thirty minutes.

Performic acid solutions are unstable, and active oxygen is lost at a fairly rapid rate (several per cent per hour at room temperature); the solutions, therefore, should not be stored but should be used immediately.

#### Epoxidation with Perbenzoic Acid

1,2-Epoxyethylbenzene (Styrene Oxide).^{12,113} To a solution of 42 g. (0.30 mole) of perbenzoic acid in 500 ml. of chloroform, prepared as described on p. 393, 30 g. (0.29 mole) of styrene is added. The solution is maintained at 0° for twenty-four hours, with frequent shaking during the first hour. At the end of twenty-four hours titration of an aliquot part of the solution shows that only the slight excess of perbenzoic acid remains. The benzoic acid is removed from the chloroform solution by shaking with several portions of 10% sodium hydroxide solution, the alkali is removed by washing with water, and the chloroform solution is dried over anhydrous sodium sulfate. Fractional distillation yields 24–26 g. (69–75%) of 1,2-epoxyethylbenzene, b.p. 101°/40 mm., as an almost colorless liquid.

cis-9,10-Epoxystearic Acid.^{3,30} To 750 ml. of an acetone solution of 0.4 mole of perbenzoic acid, prepared as described on p. 394, 85 g. (0.3 mole) of oleic acid ^{114,115,116} is added at 0-5°. The solution is allowed to stand for forty hours at room temperature and then cooled to -25° and filtered; the precipitate is washed once with cold acetone. The crude 9,10-cpoxystearic acid (purity 95-99%) is a white powder weighing about 85 g. Two recrystallizations from acetone at 0 to -25° yields 55-60 g. of analytically pure cis-9,10-epoxystearic acid, m.p. 59.5-59.8°. Oxirane oxygen: ¹¹⁷ calcd., 5.36%; found, 5.33-5.37%. The yield is

Hibbert and Burt, Org. Syntheses, 8, 102 (1928); Coll. Vol. I, 494 (1941).

Brown and Shinowara, J. Am. Chem. Soc., 59, 6 (1937).
 Swern, Knight, and Findley, Oil and Soap, 21, 1 (1944).

¹¹⁵ Wheeler and Riemenschneider, Oil and Soap. 16, 207 (1939).
117 Swern, Findley, Billen, and Seanlan, Anal. Chem., 19, 414 (1947).

1,2-Epoxy-2-methyl-3-butene (Isoprene Monoxide) (preferential oxidation of one ethylenic linkage in a conjugated diene). 118 To a stirred solution of 16 g. (0.235 mole) of isoprene in 50 ml. of ethyl chloride cooled in an ice bath a cold solution of 30 g. (0.217 mole) of perbenzoic acid in 150 ml. of ethyl chloride is added from a dropping funnel. The contents of the flask and dropping funnel are protected from moisture by drying tubes. After the perbenzoic acid solution has been added, the reaction flask is allowed to stand in a refrigerator until the oxidizing agent is completely consumed (approximately twenty-four hours). The solution is then shaken cautiously with double the calculated quantity of sodium bicarbonate solution (30 g. per 100 ml. of water) in a cooled separatory funnel until evolution of carbon dioxide ceases. The aqueous layer is discarded, and the ethyl chloride solution is dried overnight in a refrigerator with anhydrous sodium sulfate. The solution is filtered, and the filtrate is distilled through a Widmer column until unreacted isoprene begins to distill. The residual material is then fractionated twice and yields 7 g. (30-40%) of 1,2-epoxy-2-methyl-3butene (isoprene monoxide).

### Epoxidation with Monoperphthalic Acid

β- and α-Cholesteryl Oxide Acetates.²⁵ A solution of 10 g. (0.023) mole) of cholesteryl acetate, m.p. 112-114°, in 50 ml. of ether is mixed with 266 ml. of an ether solution containing 8.4 g. (0.046 mole) of monoperphthalic acid. The solution is refluxed for six hours, and the solvent is removed by distillation. The residue is dried under reduced pressure and digested with 250 ml. of chloroform which has been dried over anhydrous potassium carbonate. The mixture is filtered, yielding 6.7 g. of phthalic acid (87% recovery) and a colorless solution, from which it which the solvent is removed under reduced pressure. The residue is crystallized from 30 ml. of methanol, giving 6.0 g. (58% yield) of  $\beta$ cholesteryl oxide acetate, which on recrystallization gives the pure product, m.p.  $111-112^{\circ}$ ,  $[\alpha]_{D}^{25}-21.8^{\circ}$ . Concentration of the filtrate gives 1.55 g. (15% yield) of  $\alpha$ -cholesteryl oxide acetate. The  $\alpha$ -isomer, purified by crystallization from ethanol, has a m.p. of  $101-103^{\circ}$ ,  $[\alpha]_{D}^{25}$  — 44.6°.

### Hydroxylation with Hydrogen Peroxide-Acetic Acid

9,10-Dihydroxystearic Acid (High-Melting Isomer). A well-stirred solution consisting of 270 g. (0.898 mole) of elaidic acid (containing 94% of elaidic acid and 6% of saturated acids), 810 ml. of glacial acctic

¹¹⁸ Pummerer and Reindel, Ber., 66, 335 (1933).

acid, and 20 g. of concentrated sulfuric acid is heated to 40°, and 123 g. of 25.5% hydrogen peroxide (0.925 mole) is added dropwise over a period of fifteen minutes. The reaction is only slightly exothermic. A granular precipitate begins to form after about thirty minutes and increases in bulk as the oxidation proceeds. The total reaction time at 40° is five hours. The reaction mixture is then poured into several volumes of hot water (95-100°) and stirred well for several minutes. The mixture is cooled to room temperature and filtered, and the precipitate is washed well with cold water. The product, which weighs about 300 g. and consists of a mixture of 9,10-dihydroxystearic acid and hydroxyacetoxystearic acids, is heated at 100° for one hour with an excess of 2 N sodium hydroxide and then poured into excess hydrochloric acid, with stirring. The granular precipitate is filtered and washed free of acid. It weighs about 280 g. (93%) and consists of somewhat impure 9,10-dihydroxystearic acid, m.p. 125-128°. zation from 95% ethanol (7 ml./g.) at 0-5° yields 220 g. (78%) of pure 9,10-dihydroxystearic acid as glistening plates, m.p. 130-131°.

#### Hydroxylation with Hydrogen Peroxide-Formic Acid

9,10-Dihydroxystearic Acid (Low-Melting Isomer).4 To a wellstirred solution of 141 g. (0.5 mole) of oleic acid 114, 115, 116 in 423 ml. of 98-100% formic acid in a 1-l. three-necked flask at 25° is added during a fifteen-minute period 59 g. of 30% (100 volume) hydrogen peroxide solution (17.5 g.; 0.513 mole; 2.5% excess of hydrogen peroxide). The reaction becomes vigorously exothermic after five to ten minutes and the mixture becomes homogeneous in twenty to thirty minutes after all the hydrogen peroxide has been added. The temperature is kept at 40° with a cold-water bath at the start and a warm-water bath toward the end of the reaction. After about two hours no further consumption of peroxide is observed, and the formic acid is removed by distillation under reduced pressure (b.p. 50°/125 mm.) in a stream of carbon dioxide or nitrogen to prevent bumping. The residue in the flask, which consists of hydroxyformoxystearic acids, is heated for one hour at 100° with an excess of 3 N aqueous sodium hydroxide, and the hot, pale yellow solution is slowly poured into an excess of 3 N hydrochloric acid with stirring. The oil, which separates, is allowed to solidify, and the aqueous layer is discarded. The white solid is remelted with hot water on a steam bath and stirred well to remove residual salts and water-soluble acids. When the oil has resolidified, the aqueous layer is discarded, and the solid is broken into small pieces and air dried. This product consists of fairly pure 9,10-dihydroxystearic acid (iodine number about 2-4, neutralization equivalent 315-320), weighs about 150-155 g. (97-99%), and melts at about 92°. The small quantity of unsaturated material present can be separated readily by grinding the material and washing it by decantation with several portions of petro-leum naphtha (hexane fraction, boiling range 63-70°). 9,10-Dihydroxy-leum naphtha (hexane fraction, boiling range 63-70°).

If purified oleic acid is not available, red oil (commercial product containing about 60–75% oleic acid) may be employed. The crude 9,10-dihydroxystearic acid obtained from this material melts at about 70–75 (compared to 92° when pure oleic acid is used), and several recrystallications from 95% ethanol are required to obtain a pure product. The yield of 9,10-dihydroxystearic acid from red oil is about 50–60% of the available oleic acid. Furthermore, the 90% grade of formic acid is satisfactory, but the reaction mixture remains heterogeneous throughout. In preparations one-tenth the size described, the 25–30% hydrogen peroxide can be added in one portion. In larger preparations the addition may require thirty minutes to one hour. In preparations five addition may require thirty minutes to one hour. In preparations five to ten times the size described, it is more convenient to pour the reaction mixture into a large volume of water and then hydrolyze the washed oily layer of hydroxyformates as described.

When 90% hydrogen peroxide is employed instead of the 30% grade, the crude dihydroxystearic acid has an iodine number of 1, instead of the crude dihydroxystearic acid has an iodine number of 2.4. With the concentrated peroxide, the quantity of formic acid can be 2.4. With the concentrated peroxide, the quantity of formic acid can be 2.4. With the concentrated peroxide, the amount employed with 25–30% reduced to about one-seventh the amount employed with 25–30% hydrogen peroxide.

1,2-Tetradecane diol.² To a well-stirred mixture of 49.2 g. (0.25 mole) of 1-tetradecene, b.p.  $158-159^{\circ}/60$  mm.,  $n_D^{20}$  1.4357 (prepared by efficient fractional distillation of the 95% commercial grade), and 295 ml. cient fractional distillation of the 95% commercial grade), and 295 ml. of 98-100% formic acid at 25°, 35 g. of 25.6% hydrogen peroxide (0.263 mole; 5% excess) is added in one portion. The mixture is heated and mole; 5% excess) is added in one portion. The mixture is heated and stirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis a indistirred for about twenty-four hours at 40°, or until an analysis and analysis at 40°, or until an analysis and analysis at 40°, or until an analysis at 40°, or until an analysis and analysis at 40°, or until an analysis and analysis at 40°, or until an analysis at 40°, or until an

with hot water and allowed to resolidify. The combined water washes are extracted with ether to remove a small quantity of dissolved glycol, and the residue obtained after evaporation of the ether is combined with the main portion of glycol. The crude glycol is broken up into small pieces and air dried, yielding about 55 g. (95%) of fairly pure 1,2-tetradeeanediol, m.p. about 65°; iodine number about 4. This is recrystallized from methanol (8 ml./g.) at 0°, yielding about 40 g. (69%) of pure product, m.p. 68-68.5°.

trans-1,2-Cyclohexanediol. To a mixture of 105 g. of 98-100% formic acid and 13 g. (0.115 mole) of 30% hydrogen peroxide, 8.0 g. (0.097 mole) of cyclohexene is added. The immiscible layers are shaken together briefly; spontaneous heating occurs, and the suspension becomes homogeneous at 65-70°, where it is held for two hours on the steam bath. Most of the formic acid is removed by distillation, and the residue is heated on the steam bath for forty-five minutes with 50 ml. of 20% sodium hydroxide. After cooling, the yellow solution is neutralized with hydrochloric acid and evaporated to dryness under vacuum. The resulting solid is distilled, yielding 10.25 g. of a fraction, b.p. 128-132°/15 mm., which solidifies immediately. Recrystallization from acetone gives 7.9 g. (70%) of trans-1,2-cyclohexanediol, m.p. 102-103°. A larger scale oxidation of cyclohexene is described in Organic Syntheses.

# Hydroxylation with Performic Acid

2,3-Dihydroxynonanoic Acid.55 Twenty grams (0.13 mole) of 2nonenoic acid is added slowly to a well-stirred solution of performic acid prepared by the reaction of 69 g. of 98-100% formic acid, 19 g. (0.5 mole) of 90% hydrogen peroxide, and 0.50 g. of concentrated sulfurie acid. The emulsified mixture is heated to 55-60° to start the reaction and is then held at this temperature for two hours while stirring is continued. The temperature is then allowed to rise to 95° until the spontaneous reaction is over (twenty-five minutes) and the excess peracid largely destroyed. Most of the formie acid is removed by vaeuum distillation, and the residue is saponified on the steam bath for onehalf hour with 175 ml. of 10% sodium hydroxide. After acidification with hydrochlorie acid, the oily product is extracted with ether and the extract is dried over anhydrous sodium sulfate. Evaporation of the ether yields a waxy solid which is suspended in benzene and filtered, yielding 2,3-dihydroxynonauoic acid as white slippery flakes. Concentration of the filtrate followed by addition of ligroin gives two additional erops,

in Roebuck and Adkins, Org. Syntheres, 28, 35 (1948).

the total yield of product being 12.4 g. (51%). On crystallization from ethyl acetate or water, pure 2,3-dihydroxynonanoic acid, m.p. 118-118.5°, is obtained.

# TABLE OF ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

The following table lists the ethylenic compounds which have been epoxidized or hydroxylated with organic peracids. The table is divided into the following sections: A, Hydrocarbons and substituted hydrointo the following sections: A, Hydrocarbons and substituted hydrointo the following sections: A, Hydrocarbons and substituted hydrointo the following sections: C, Acids; D, Alcohols; E, carbons; B, Steroids (alphabetical order); C, Acids; D, Alcohols; E, Esters; F, Aldehydes and ketones (including carbohydrates); G, Ethers; H, Miscellaneous.

In the preparation of the table the literature has been consulted to October 1951. The addendum to Table I lists the compounds whose epoxidation or hydroxylation with organic peracids was reported from October 1, 1951, to October 1, 1952.

TABLE I

ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

	ETHYLEN	ETHYLENIC COMPOSITE					
	button of the state of	Yield of	Yield of Oximne, % (Reference)	reneo)	Yield of	Yield of α-Glycol, % (Reference)	ereneo)
	Ethylenia Compound						
Formula	Мато	Perbenzoie Aeid	Monoper- phthalic Aeid	Peracetic Acid	Peraeetie Acid Performie Aeid	Performie Aeid	Perbenzoic Aoid
		A. Hydrocarbons and Substituted Hydrocarbons	and Substituted H	ydrocarbons			
						<u> </u>	
7117	Ethylene-C14	30-53 (120)				73 (122)	
110	1,3-Butadiene	42 (118, 121)				Low (123)	
Callabra	1,4-Dibromo-2-butene					30 (123)	
CAHCH	1,4-Dichloro-2-butene					<u> </u>	
•	3,4-Dichloro-I-butene				70 (124)		
CILIC	3.Chloro-2-methyl-1-propene						
	(methallyl chloride)					(122)	
C,II,s	1-Butono				54 (71)	85 (122)	
5	1 Chlore Lenglopontone	75-80 (125, 126)					
Carro	1-Chloro-2-cyclopentene	75-80 (125, 126)					
Colls		80-90 (70, 127)			(129)		
	Isoprene	30-60 (118, 121,					
	3-Methyl-1.2-butadiene	(02)			(129)		
	1,4-Pentadiene	— (121)		(00)			
CsH10	Amylenes			(23)			
Cell'1N	-cyclopentene	(130)					
CeIIs		27 90 (131)					
Colloci	1-Chloro-1-eyclohexene	132, 133)					
	1-Chloro-2-cyclohcxene	75-80 (125, 126,					
		(101					

30 (142)						
65-75 (55, 119, 122, 141)	58 (141, 144)	70 (1:14)		59 (141) 30 (141) 73 (141)	81 (141) 40 (141)	
63-100 (32, 70, 138, 139, 1:10)		(147)		(140)		— (129) — (129)
60-67 (32, 137)			(149)			
				.6.	04)	
— (135) 100 (70, 136)	— (121) 66 (121, 143) — (121) 75 (10, 70, 144) — (132) — (132)	85 (145, 146) — (7)	93 (148) — (130) — (120) 75-80 (125, 126,	134) — (150, 151) 100 (152) — (121) 50-75 (10, 70, 136,	153) 55 (132, 136, 154) 60-90 (136, 155) — (132) — (150) 70 (125)	31 (157) — (158) — (159) — (159)
Biallyl 100 Cyclohexene		jo	-huteno a 1-4-ono syclohexene cyclohexene	<b>6</b> 0		
C ₆ H ₁₀	8-8-8	C ₆ H ₁₁ BrO C ₆ H ₁₂	C,H;0 C;H;0 C;H;0 C;H;0	C ₇ H ₁₂	ć t	07H120 07H1301 07H14

TABLE I-Continued

# ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

	ETHYLEN	ETHYLENIC COMPOUNDS UNIDIZED WITH CHARGO	OXIDIZED WIT	A Charles			
	Estrabatic Compound	Yield of	Yield of Oximne, % (Reference)	renee)	Yield of	Yield of a-Glycol, % (Reference)	erenec)
Formula	Мато	Perbenzoic Acid	Monoper- phthalic Acid	Peracetic Acid	Peracetia Acid	Performie Aeid	Perbenzoio Aeid
		trademakan and Substituted Hydrocarbons-Continued	ubstituted Hydroco	rbons—Continued			
Calle	Cyclodetatetraene Styrene	40-60 (160, 100a) 69-75 (12, 22, 113,		55 (101)		40 (122)	
C.II.Br.Cl.	7.8-Dichlorobioyelo-[4.2.0]-dibromo-	— (160)			-		
	octeno	****					
C, II, Cl	7,8-Dichlorobicyclo-[4.2.0]-2,4-octadiene — (150) 1-Vinylcyclohexone — (121)	- (121) - (121)			(165)		
: ;	4.Vinyleyelobexene	65 (160)		69-80 (165a)			
<b>.</b>	l-1-cyclohexene	75 (166)				•	
-	1,3-Dimethyl-I-cyclohexene 2,4-Dimethyl-I-cyclohexene	— (132) — (132)					
	Dimethyleyelohexene	6					
	2,5-Dimethyl-1,5-hexadione	- (121) - (133)					
	3-Methyl-1-methylenecyclonexund	(132)					
	1-n-riopy-1-cyclopeaces	— (132)					
	1,7-Octadiene	(121)					
C ₆ H ₁ (O	1-Ethoxy-1-eyelohexene	70 (125)					
-	3-Ethoxy-1-cyclohoxene	<b>—</b> (155, 167)					
$C_6H_{13}C1$	2-Chloro-2-oeteno	25 (157)			(41, 43, 168)		
$C_8II_{16}$	Diisobutylene 2-Methyl-1-heptene	(e. c)			(129)		_

		EPO:	XIDA	TION		OK E	11	11	111	911	10										
													(183)								
58-70 (2, 169)	— (36) 40 (40)																				
	1, 147)	100 (138)	100 (138)						(168)			(100 (139)									-
(35 (2)	28 (21) (36, 40) (36, 40)																				
_					(178)											5, 186)		174)		, 174)	-
	15 (21) 40 (36) 70 (40)	25 (170) 100 (70, 132, 163, 171)	(172) 80 (172) (170)	60-80 (173, 174) - (162, 175, 176) 80 (162, 177)		- (132) - (132, 179, 180) - (121)	(132)	(132)	100 (181)	70 (127, 171)	(182)	-(163, 182)	-(121)	75 (172)	75 (172)	ne (75-80 (125)   100 (184, 185, 186)	(175)	60-80 (173, 174)	(136)	60-80 (173, 174)	-
		rinethyl-z-pentene thoxy-2-butene				oleyclohexene ylidenecyclohexane	1,8-Nonadiene		1-Nonene	Icononene	1.2-Dihydronaphthalene	2,3-Dihydronaphthalene	Divinylbenzene	cis-1-Phenyl-1,3-butadiene	4-(p-Bromophenyl)-1-butene	2-Chloro-1,2,3,4-tetrahydronaphthalene 75	1-Phenyl-J-butene	4-Phenyl-1-butene	1,2,3,4-Tetrahydronaphthaleno	1.Anisyl-1-propens 5-Phenyl-1-pentene	
	1-Octen   Octens   2,4,4-Tr	C ₈ II ₁₆ O ₂ 1,1-Die C ₉ II ₈ Indene		CoHoCi 3-Ci CoHio Ally		Collie 16	==		Collin	: :	Clattia			0.11.13.	Cigniliza	ClelfiiCl	C101112			CleH12O	

TABLE I-Continued

RGANIC LEKACIDS
5
WITH
OXIDIZED
COMPOUNDS (
ETHYLENIC

	LTHYLEN	ETHYLENIC COMPOUNDS OXIDIOED WITH	OALDIAN WAS				
	Ethylenic Compound	Yield of	Yield of Oxirane, % (Reference)	rence)	Yield of	Yield of a-Glycol, % (Reference)	ereneo)
Formula	Name	Perbenzoie Aeid	Monoper- phthalic Aoid	Peracctic Acid	Peracetia Acid	Peracetia Acid Performic Acid	Perbenzoic Acid
	A.	A. Hydrocarbons and Substituted Hydrocarbons-Continued	ibstituted Hydroca	rbons—Continued			
CroHis	Сатрвене	- (187)		(188)	(189)		
	(+)-A'-Carene (+)-A'-Carene	70 (31)		69 (31, 188)	(31)		
	2,4-Dimethyl-4-vinyl-1-cyclohexene Limonene	- (190) 40-60 (6, 32, 101, 191, 192)		63 (32)	— (193)		
	Myrcene	)		25 (194)	— (195)		
	Norbornylene Pinene	— (6, 187, 191, 196, 197)		89 (31, 198)			(142)
	Sabinene a-Terpinene	(199)					
$C_{10}H_{18}$	γ-Terpinene 1-Butyl-1-cyclohexene	40 (200) — (132)					
	1,9-Decadiene 2,6-Dimethyl-2,6-octadiene	- (121) (121)		1000	(4)		
	3-Monthene	83–91 (201) — (132)		(102) 08-60	(011)		
C10H20	Caprylene 1-Decene	— (6, 191) 100 (181)		56 (2)		45-75 (2)	
C11H10 C11H100	Decene 1-Phenyl-3-penten-1-yne 1-Anisyl-1-butene	(- (6) (- (175)		50 (34)			

	EP	OXI	OITAGI	N	)F E	тнх	יננ	JK10								
											6	40-75 (2, 212)				-
												- (213)		40 (58)		
					62 (34)							52 (2)				
•																
— (10, 70, 202) 90 (203a) 60-80 (203, 204)	60-80 (176, 203) — (205)	— (175) 70-90 (184)	— (175) — (185, 186, 206) 70–80 (207)	70-80 (201)	— (132) 100 (181)	— (164) 100 (10, 70, 208,	209)	- (202) - (203a) - 70-90 (184)	(70-90 (181)	(210)	(100)	100 (181, 211)	416/	:   	(203)	(
1-Phenyl-1-cyclopentene 90 3-Phenyl-1-cyclopentene 60.			-butene -butone -propene yl)-2-methyl-1-		<del></del>	oxen-1-yne ystyreno	1-Phonyl-1-eyclohexene	1-Aninyl-1-eyclopentene 3-Aninyl-1-cyclopentene	1-Phenyl-2-ethyl-1-butene   1-Phenyl-2-methyl-1-pentene	6.Phenyl-1-hexene   4.Anisyl-2-methyl-1-butene   1.Anisyl-1-pentene	3-Anisyl-2-pentene 1,2,5-Trimethyl-5-isopropenyl-1-cyclo-	hexeno	Indoderens	3-Ethoxy-4-propyl-3-heptene	1-Phenyl-3-methyl-1-cyclohexene	1.Benzyl-1-cyclohexene
	Cullia03   I-C		Cullio 1-			Cullin Cullin	Challin	C111110	CreHis	Cullio	Collec	0711713	Cirilra	C11[[10]	Chillis	Cully

TABLE I-Continued

ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

	ETHYLEN	ETHYLENIC COMPONENCE					
	Ethelenia Campound	Yield of	Yield of Oxirnna, % (Reforence)	srence)	Yield of	Yield of a-Glysol, % (Reference)	sronce)
			Monoper-	Porgonia Aold	Peracetic Acid Performic Acid	Performie Acid	Perbenzoio Acid
Formula	Namo	Perbenzoio Acid	Acid				
		A. Hydrocarbons and Substituted Hydrocarbons-Continued	ubstituted Hydroco	rbons—Continued			
	-						
Ctalliso	1-Aninyi-2-othyi-1-buteno	(266)					
C, 11, 9C)	1-Tridecone trans-1,1'-Dichlorostilbene	50 (215)			(216)		
C141113		- (217)		<b>—</b> (137)			
		60-100 (22, 185, 216, 217)		83-100 (137, 201)	(612)		13 (218)
	_		(010)				(2)
CidIIs	1-Phonyl-2-oyelohexylethylene		(617)			70-80 (220)	
Citilio	1-(2,3-Dimethexyphenyl)-1-cyclenexene	- (210)					
Citilia	ylone	76 (221)		(000 0/ 01		49-95 (2, 223)	
C141133	1-Tetrndeceno	***************************************		(777 '7) 21.			
C18111202	1.Phonyl-2-(3,4-methylenedioxy-	— (203, 204)					
:	phonyl)cthylene	- (100, 224, 225)					
C161114	1,1-Diphenyt-1-propend	(220)					
	_	— (175)			1961, 000		
					(961) 001		
	/l) ethylene	(176, 203, 227)					
CisHiO	1.Phenyl-2-anisylethylene	(223)					
	1-Phenyl-1-(o-methoxyphenyl)ethylene	(220)					_

# TABLE I—Continued

ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

						100 (Bof	(worder
	2	Yield of	Yield of Oxirane, % (Reference)	reneo)	Yield of	Yield of a-Giyeoi, 70 (restoration)	
	Ethylenie Compound						
Formula	Name	Perbenzoio Acid	Monoper- phthalic Acid	Peracetie Acid		Peracetic Acid Performic Acid	Perhenzoic Aeid
		A Hidrocarbons and Substituted Hydrocarbons-Continued	Substituted Hydroc	arbons—Continued			
C24H24	enylyl)-1-phenyl-2-etbyl-1-	75 (242)					
C24H32 C26H20	buttan 1,1-Diphenyldodecone Tetraphonylethylene	— (243) 100 (244) — (245)					
C361148		(245, 246) (247)					
CroHso	l hydroearhon 6-amyrilene	(248) - (246) - (22)					
# 5	Squalene 2-Lupene a-Carotene		70 (249) — (250)				
O401156	β-Carotene Carotene (mixture of isomers)	10-15 (252)	(207)				
C42H32 (C5H8)n	1,2-Bis(benzyl-9-fluorenyl)ethylene Rubber	65 (253) — (254, 255)		— (256)	(256, 257)		
		B. Stero	B. Steroids (alphabetical order)	rder)			
	38-Acetexyallopregnan-20-one enol acetato	(258)					

TABLE I—Continued

ERACIDS	Yield of a-Glycol, % (Red
ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC FERACIDS	) Xield

Yield of a-Glycol, % (Reference)		Peracetic Acid Feriorano Acid	-										_			- (304)		(304)	20 (304)	_
χ̈		Peracetic 4		- (293)		— (293)			25-30 (298)				_							
rencc)		Peracetic Acid	Continued									30 (300)								
Yield of Oxirane, % (Reference)	Monoper	phthalio Acid	B. Steroids (alphabetical order)—Continued		25-60 (294, 295)	ì				40 (299)		70 (050 301)	(*00, 600)							
Yield of		Perbensoio Acid	B. Steroids (alp		50 (294)		(100)	(288)	Fair (296)	40 (297)		20 (300)	25 (301)	80 (302)		70 (303)	(305)	60 (306)		
	Ethylenia Compouna	Name				trans-Dehydroandrosterone henzoate	tetracetylglucoside	Dehydroergosteryl acetate-malcie	anhydride adduct Debydrojsoandrosterone	Dehydroisoandrosterone acetate	3,6-Diacetoxy-5-methyl-10-norangrost	<b>6</b> 1		pregnadione	Diarotacharia adduct	3,7-Dihydroxycholenie acid	Dibydroergosteryl acetate	3a,12a-Dihydroxy-14-choleme acid	a-Ergostenyl acetate	Theresia
		Formula				·		-												

	80 (269) — (317)	100 (320)	80 (203)
- (302) - (303) - (303) 25 (300) 80 (310) - (258) 71 (315) 56 (311)	- (312) 70 (313) - (314) - (310) 60-70 (201, 288,	118) 50 (319) 10 (319) - (259, 307) - (259, 307) 10 (- (259) 110 (- (259)	— (323) — (318, 324) — (274)
acetate-maleic anhydrido nol-3a-one-17 ion-3a-ol-17-ono acetate y-11-cholenie acid ypregnan-20-ono enol acetate lonpirostene soallospirostene-36-0-3-	Actiato Inciliadroycholenio acid 3.Ketoandrotta-4,16-dieno a.Methoxy-16-i-pregnen-20-0no A.Methyl 3g-acetoxy-14,10-alloetiochola- dienato Methyl 3g-acetoxyallo-14-etiocholenate Methyl 3g-acetoxyallo-14-etiocholenate Methyl 3g-acetoxyallo-14-etiocholenate Activityl 3g-acetoxy-6,11,10-chola- friento	Methyl 3a-nectury-11-cholenato Nethyl 3g-nectory-11-cholenate Methyl 3g-nectory-14,19-etionllochola- dienato Methyl 3g-nectory-14,19-etiochola- dienato Methyl 3g-nectory-3(11)-etiocholanato Methyl 3g-nectory-3(11)-etiocholanato Methyl 3g-nectory-3g-etionllocholenato Methyl 3g-nectory-12g-bydroxy-7-	cholenate Methyl 3f-acetoxy-14,17-isoalloctio- cholenate Methyl 9-cholenate Methyl 11-cholenate Methyl 7,14-3c,12f-diacetoxychola- dienate

TABLE 1—Continued

FTHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

	Campound	Yield of	Yield of Oxirane, % (Reference)	renco)	Yield of	Yield of a-Glycol, % (Reference)	erence)
Formula	Nanie	Perbenzoie Acid	Monoper- pluthalic Acid	Peracetic Acid	Peracetio Acid   Performic Acid	Performie Acid	Perbenzoic Acid
		B. Steroids (al)	B. Steroids (alphabetical order)—Continued	Continued			
	Methyl 3a,12a-dinectoxy-14-cholenate Methyl 3a,12g-dinectoxy-14-chelenate Methyl 3a,12g-dinectoxy-14-chelenate Methyl 3a,12g-dinectypapochalate Methyl 3a-fallydroxy-14-chelenate Methyl 3a-dilydroxy-11-chelenate Methyl 3a-hydroxy-11-chelenate Methyl 3a-hydroxy-11-chelenate Methyl 3c-hydroxy-12-methoxy-9,11- chelenate Methyl 3-keto-1,11-cheladicate Methyl 3-keto-1,11-cheladicate Methyl 3-keto-1,11-cheladicate Methyl 12-methoxy-9,11-chelenate 5-Methylmorcholestane-3,6-dilone 3-Methyl 12-methoxy-9,11-chelenate 5-Methylmorcholestane-3,6-dilone 3-Methylmorcholestane-3,6-dilone a-Methylmorcholestane-3,6-dilone 3-Methylmorcholestane-3,6-dilone a-Methylmorcholestane-3,6-dilone 3-Pregnadien-3-one noctate Pregnanonel acetate 17-Yinyl-3,17-iscandrostanediol	- (250) 50 (274) - (312) - (312) - (374) 45 (310) 55 (310) 56 (325) - (325) - (325) - (325) 70 (207)	>40 (327) (328) (330) 05 (263, 331)	55 (300)	25 (208)		. (272)

TABLE I—Continued

ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC PERACIDS

erence)		Perbenzoie Acid					(171)	(171)						
Tr. 11 - Clumbl % (Reference)	ardigues of	Peracetic Acid Performic Acid		69 (350) 88 (350) 73-94 (350,	3664, 367) 80-94 (349, 350,	3873) 68 (350) 60 (350)	(20)			57 (373)			- (375)	
7	Y left of						20 (368) — (369)	(369)	(320)	40 (371, 372)	- (344) - (338)	— (357) 58 (338, 357)		
	rence)	Peracetic Acid								— (370)				
	Yield of Oxirane, % (Reference)	Monoper- phthalio Acid	C. Acids-Continued	1	(300 <i>a</i> )								( (374)	
	Yield of	Perbenzoic Acid	G. A									55 (357) 70 (171, 357)		
	Ethylenia Compound	Name		cis-10-Octadecencie	cis-11-Octadecenoio	trans-11-Octadecenoic	cis-12-Octadecenoio trans-12-Octadecenoio Vaccenic	cis-12-Hydroxy-9-octadecenoid	(nemotec.) trans-12-Hydroxy-9-octadecenoid (ricinelaidic)	α-9-Octadecene-1,18-dicarboxylic n-11-Eicosenoic	Anacardic 9,10-Diacetoxy-12-octadecenoic		Erucic       Elemolio  Minod uncontrated fatty acids from	human hair fat
		Formula		C18HMO2	(cont a)			C18H34O3		C20H36O4 C20H38O2	C22H32O3	C22H4004 C22H4202	C30H48O3	C12H42O2

# TABLE 1—Continued

austradjus idaser seminadistrurridingenin	(erence)	Perbenzoic Aeid								
entripe o de plantague adague e tra de la company	Yield of a-Glycol, % (Reference)	Peracetia Acid   Performio Acid		(378)		50 (55)		30 (401)	20 (401)	— (55) —
ERACIDS	Yield o	Peracetic Acid							— (105) — (405)	— (405) — (405)
rii Organic P	(erence)	Peracetic Acid				32 (402)			10 (1, 23)	
Oxidized we	Yield of Oxirane, 55 (Reference)	Monopor- plethatio Acid	E. Estera			>32 (102)	65 (103) 51 (402) 60 (403) 60 (335)			
Erayleng Compounds Oxidized with Organic Peracids	Yiebl o	Perbenseis Acid		- (303, 309) - (109)	— (125) — (101) — (125, 180) — (121)	70 (377)			— (387) — (121) — (387)	
	1 hyleste Companie	Natur	Managaman der des anders der	Vnyl neetato Methyl 2, Eberadianouto (sorbato) Ethyl neetsacotato	1.Aretoxy-1-gyelohereno 1.Aretoxy-3-methyl-l-cyelohereno 2.Aretoxy-1-methyl-1-cyelohereno Methyl diedlyheretato	Diethyl allylinalonata Methyl 2-nonenoate Cinnanyl acetato	Methyl ceallogyclogoranato Diethyl (1-methyl-2-propenyl)malonato Ethyl ceyclogeranato Methyl 11,3-trinothyl-3-cyclohevene-2-	Ethyl 5-cyclopentyl-5-hydroxy-2-	ite (undecylenate) ie 5 (undecylenate) idecenoate	Dimethyl traunatate Propyl 10-hendecenoate (undecylenate) 2-Methoxyethyl 10-hendecenoate (undecylenate)
	, , , , , , , , , , , , , , , , , , ,	P. gradie		Calko; Calked; Calked;		Challao		C12111301		Chillings   12   12   13   14   15   15   15   15   15   15   15

	EPO	OITAGIZ(	N OF I	ETHYL	ENIC	COMP(	OUNDS		419
T									
	6 (37.• 50)	50 (56, 40%) 100 (1, 169)							
— (49) <20 (315)	50 (49, 72, 407)   96 (37, 56) (49, 407) (409)		(36)	— (29, 51, 361, 362)		Good (357) — (357)			
- (28, 20, 30)	45 (1, 23, 355) 5(-72)	- (1, 23)		40 (410)	(355)		(413)	— (118a)	
	÷ 1.				_	- (413)	(374)	(418)	
0 00 345.	25-10 (23, 23, 23, 24, 24, 200) 20 (29, 310) 42-67 (3, 20) — (171) S0 (408)	— (408) 80 (29)	85-95 (28, 20)	— (343) — (171, 240, 342.	(240)	- (411) - (411) - (357) 75 (357)	— (247) — (414) — (395)	(416, 417)	(247)
	ate)		Methyl toract	Subject of the state of the sta	Ethyl trans-9-octadecenoate	Olcyl acetate Methyl (+)-pimarate Methyl (+)-dihydropimarate Methyl brassidate   Methyl erucate	Octyl oleate Methyl a-elemolate A-Amyrin acetate Buphadicnyl acetate Euphadicnyl acetate	Euphol acetate Germanicol acetate Lanosteryl acetate	Artenyl acetate Euphenyl acetate
	17H2202 Methyl palm 18H3402 Methyl 9.12- (linoleath) Methyl 0.11- Methyl 0.13- Methyl 0.13- Methyl 0.13- Methyl 0.13- Methyl 0.13- Methyl 0.13-	Methyl cis- selinate) Methyl tra selinate) Methyl tra Sclaidate			Czonsacz	C21H32O2 C21H33O2 C23H44O2	C26H50O2 C31H50O3 C32H52O2		C32H54O2

C19H36O2

C₁₇H₃₂O₂ C₁₉H₃₄O₂

TABLE I—Continued IPOUNDS OXIDIZED WITH ORGANIC PERACIDS

	rence)	,	Perbenzoic Acid									
	Yield of a-Glycol, % (Reference)		Performic Acid			— (37)						
	Yield of		Peracetic Acid Performic Acid				•				36 (52, 361, 362, 369, 428)	— (50) — (428)
a Ondaria	rence)		Peracetic Acid		— (355)		— (355)		86 (1, 427)	(355)	73 (1, 427)	70-80 (1, 427, 429)
OXIDIZED WITH	Yield of Oxirane, % (Reference)		Monoper- phthalic Acid	E Release-Continued		— (422)	— (423) — (424)	6 (425) — (425)	— (426)			
ETHYLENIC COMPOUNDS OXIDIZED WITH ORGANIC	Vield of		Perbenzoie Acid	- 2	— (247) 70 (420) — (247) — (421)	— (418 <i>a</i> )						
ETHYLEN		Ethylenic Compound	Мате		Euphorbenyl acetate Isodiiydrocuphol acetato Trivoslienyl acetato Methyl acetyleburicoate	Oloyl olento Artenyl benzonto Beningenin tetrnacetate	Propylene glycol dioleate Isoescingenin pentracetate Diethylenerlycol dioleate	Cryptoxantliin diacetate Xanthophyll diacetate	Capsanthin discetate Triolein	Butyl Carbitol esters of unsaturated	fatty acids Castor oil	Cocoa butter Coconut oil Corn oil
			Formula		C32H54O3 (Cont'd)	Cselfs:02 Csrlfs:02	C10H12O10 C10H12O10 C10H14O6	C44H6003	C. Hr2Os	2010110		

EPOXIDATION OF E	THY	LENIC COMPOUNDS 421
		75 (438, 439)  — (410)  — (438, 441) 30 (412) 33 (440) 30 (441, 442) — (444) 90 (439)
— (430) — (37,* 434)		
- (428) - (419) - (431-436) - (358, 437) - (50)	- E	
71 (1, 427) 74 (1, 427) 66 (1, 427) 57 (1, 427) — (355) 77 (1, 427) 75 (1, 427) 76 (1, 427) 71 (1, 427) 71 (1, 427) 71 (1, 427) 71 (1, 427) 71 (1, 427) 73 (1, 427)	okydrale derivativ	
	es (including earb	. (446) 60-70 (446) 50 (447)
	R Aldehudes and Retones (including carbohydrale derivations)	- (438, 439) - (3) - (3) - (4) - (6, 9) 59 (443) - (6, 8) 96.5 (445) 86 (445)
Cottonseed oil Linsect oil Linsect oil Merhaden oil Methyl esters of soybean oil acids Methyl esters of unsaturated acids Neatsfoot oil Olive oil Perila oil Rapeseed oil Ripeseed oil Ripeseed oil Sardine oil Sybean oil Tall oil Tall oil Tall oil Tall oil		Rhamnal Galactal Galactal Gucal 3-Methylglucal Methylheptenone Benzylideneacetone Citral Pulegone Citral Triacetylgulactal Triacetylgulactal Lactal Caclobial ca-Ionone p-Ionone
		C ₆ H ₁₀ O ₃ C ₆ H ₁₀ O ₄ C ₇ H ₁₂ O ₄ C ₈ H ₁₄ O C ₁₀ H ₁₀ O C ₁₂ H ₁₀ O C ₁₂ H ₁₀ O C ₁₂ H ₁₀ O C ₁₃ H ₁₂ O

^{*} Oxirane formed.

TABLE I—Continued

FERACIDS	
ORGANIC	
WITH	
OXIDIZED	
COMPOUNDS OXIDIZED WITH ORGANIC FEMALES	COMPONIO
ŗ	THATENIC THATENIC

	NATARA	ETHYLENIC COME COLL					
	-	Yield of	Yield of Oxirane, % (Reference)	srence)	Yield of	Yield of α-Glyeol, % (Reference)	crenee)
	Ethylenie Compound						
Formula	Name	Perbenzoie Acid	Monoper- phthalic Aeid	Peracetic Aeid	Peracetic Acid	Performie Aeid	Perbenzoie Aeid
				deringtines) -Cor	linued		
	F. Aldehy	F. Aldehydes and Ketones (including carbonyurate at the state of the s	luding carbonyara	( a m )			
	11-Keto-1-trideeene	- (387)					30 (448)
C13H240 C14H150 C14H220	Tetracety-1-glucosene Methyl a-fonone 11-Keto-1-tetradecene	45-55 (387)	60 (445)				
C14120 C161240 C171240 C101140	a-Dilydroionone ethylene ketal 11-Keto-11-phenyl-1-hendeene Lanostenone	— (387) — (416) — (247)	(dr.) 00				
Cyonsoc	Tribitorione						
			G. Einers				
							(450)
C'H'O	Furan Ethyl vinyl ether	25 (451)					
C,H ₃ O	yran	58 (452) $- (121)$				71 (453)	
C6H1002		45 (170)					
CoH100	Phenyl allyl ether	35 (454)			100 (138)	25-33 (455)	
C10H10O2	Isosafrole	(203)			100 (155) — (138)		(22)
C.H.	Safrole Anethole			62 (24)	55-100 (32, 138)		(450)
\211mO	Methyl einnamyl ether	85 (377, 457)			100 (138)		
$C_{10}H_{12}O_{2}$	Eugenol	_		-			

Ţ	EPON	IDATION OF ETHYLENIC COMPOUNDS	420
95 (459)		— (370)	
		(460) (460) (460) (460)	
. (32) –			— (355) — (355)
	H. Miscellancous	— (461) 69 (461) 85 (461)	
— (457) 50 (454) 25 (454) 60 (458)	H.	8 (462) 30 (463) - (464) 60 (463) - (339) - (30)	r.
Isocugenol Ethyl cinnamyl ether Allyl einnamyl ether Sallyl ether Dispiro[dicyclohexane-2,5-dihydrofuran] Gardanol methyl ether		Butadiene sulfone  2-Ethyl-2-pertennamide  2-Ethyl-2-pertennamide  2-Ethyl-2-pertennamide  2-Ethyl-2-pertennamide  3-Ethyl-2-pertennamide  Furfural diacetate Benzyl propenyl sulfone  Furfural phenyllydrazone  Thiopyrine  Benzaldehydephenylhydrazone  V-Santonin  Oleamide  N-Aettyloleamide  N-Aettyloleamide  N-C-Hydroxyethyl)oleamide  N-C-Hydroxyethyl)oleamide  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Hexyloleamide)  N-(n-Mexyloleamide)  N-(n-Mexyloleamide)	
C1H140	-	C4H602S C6H602S C6H602S C6H1002S C7H11000 C9H11002 C9H11002 C1H11002S C1H11002 C1H11002 C1H11000 C2H11100	Carlino

# ADDENDUM TO TABLE I

The compounds appearing in this addendum are listed alphabetically in sections which correspond to those in Table I.

Compound	Peracid	Product	Yield	Reference
A. Hydroca	arbons and Substituted	Hydrocarbons		
-Acetoxy-1-cyclohexenc	Peracetic, performic	Triol	20-25	468
	Peracetic	Oxirane	****	469
	Monoperphthalic	Oxirane		470
	Performic	Glycol	42	471
	Performic	Glycol	42	471
	Peracetic	Glycol	30	468
-Phenyl-1-(2-biphenylyl)ethylene	Perbenzoie	Aldchyde (vin the oxirane)	<del>-</del>	470
·	B. Steroids			
3β-Acetoxy-7,8-epoxy-9(11),22-ergo- stadiene dibromide	Perbenzoic	Glycol	_	472
38-Acetoxy-7,9(11),20-ergostatriene	Perbenzoic	Oxirane	-	472
3β-Acetoxy-7,9,22-ergostatriene	Monoperphthalic	Oxirane	l —	473
16,20(22)-Allofurostadiene-3\$,26-di- ol diacetate	Monoperphthalie	Oxirane	_	474
Allopregnane-11,20-dienol acetate	Perbenzoie	Glycol	-	475
8(14)-Androsten-3β,17β-diol diace- tate	Monoperphthalie	Oxirane	10-35	476
9-Androsten-3a-ol-17-one	Perbenzoic	Oxiranc		477
3β-Benzoxy-7,9(11)-cholestadiene 3β-Benzoxy-7-cholestene	Monoperplithalic	Oxirane	70	478 478
2-Cholesten-6-one	Monoperphthalic	Oxirane	50	479
3\$,17\$-Diacetoxy-7,9(11)-andro-	Perbenzoie	Oxiranc	40	478
stadiene 22,23-Dibromo-3β-acetoxy-	Monoperphthalic Peracetic	Oxirane Oxirane	40	472
7,9(11)-ergostadiene		Oxuane		
7,9(11),22-Ergostatrien-38-ol acetate	Perbenzoic	Oxirane	l _	480
9-Etiocholen-3α-ol-17-one	Perbenzoic	Oxirane		477
Methyl 3α-acetoxy-7,9-choladienate	Monoperphthalic	Oxirane	_	473
Methyl 3α-hydroxy-9(11)-cholenate	Perbenzoic	Oxirane	-	481
5\$-Methyl-3\$-methoxy-19-nor- coprost-9(10)-en-6-one	Peracetic	Oxirane	-	482
5β-Methyl-19-norcoprost-9(10)-en- 3β,6β-diol diacetate 9(11),17(20)-Pregnadiene-	Peracetic	Oxirane	-	482
3a,11,20-triol trincetate 9(11)-Tigogenin acetate	Perbenzoic	Oxirane	-	483
	Perbenzoic	Oxirane	_	481
	C. Acids			
cis-9-Hendecenoie	Performic	61	1 00	404
trans-9-Hendecenoic	Performio	Glycol	30	484 484
	1	Glycol	55	454

# ADDENDUM TO TABLE I—Continued

Compound	Peracid	Product	Yield	Reference
	E. Esters			
		Oxirane	20	469
-Amvrin acetate	Peracetic	Oxirane	50	469
-Amyrin benzoate	Peracetic	Tetraacetate	57	485
is 9 Puter 1 4 diel discetate	Peracetio	_	51-79	485
rans-2-Buten-1,4-diol diacetate	Peracetic, performic	Tetraacetate, formates	0	486
	Perbenzoic	Oxirane	_	487
Methyl acetylebullcoate	Perbenzoic, peracetic	Oxirane	80	
atomy, morotate access		Oxirane	_	487
Methyl morolate benzoate	Peracetic	Oxirane	_ '	487
Moradiol diacetate	Peracetic	Oxirane	_	488
x-Noramyrenonyl acetate	Perbenzoic	Not isolated	_	489
Peach oil	Peracetic	Oxirane	_	490
Zeorinin acetate	Peracetic	Oxirane	_	490
Zeorinin benzoate	Peracetic	O'ALLEAN !		
		Glycol	50	491
Butyl p-(2-methylalloxy)benzoate	Peracetic	Glycol	_	491
m-Carbobutoxyphenyl 2-methallyl	Peracetic	0.300.		
ether		Glycol	50	491
4-Chloro-3-methylphenyl 2-methallyl	Peracetic	Gi, year	Ì	491
ether		Glycol + oxirane		491
p-Chloropbenyl 2-metballyl ether	Peracetic	Glycol	6-50	491
3,5-Dimetbylphenyl 2-methallyl etber	Peracetic, performic	Glycol	33	
2-Methallyl m-nitrophenyl ether	Peracetic	Glycol + oxirane	42 + 25	491
2-Methallyl phenyl ether	Peracetic	Glycol	6-25	491
2-Methallyl m-tolyl ether	Peracetic, performic	Glycol + oxirane	20	491
2-Methallyl o-tolyl ether	Peracetic	Glycol + oxirane	60	492
2-Methallyl p-tolyl ether	Peracetic Performic	Glycol	25	492
5.6-Dihydro-2-pyran	Performic	Oxirane	25	1
2,5-Dihydro-2,2,5,5-tetramethylfuran	Performic		<u> </u>	
	H. Miscellaneous			
	II. ALL EGOOGTA		1	1
		1	1	491

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